MEETING ABSTRACTS

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PHYSICIAN ABSTRACTS

Oral communications

O1 Pulmonary embolism related sudden cardiac arrest admitted alive at hospital: characteristics and outcomes

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Introduction Pulmonary embolism (PE) is a relatively common cardiovascular condition, occasionally and tragically manifesting as sudden cardiac arrest (SCA). The natural history of SCA complicating PE has been poorly evaluated. Guidelines suggest the consideration of thrombolytic therapy when PE-related SCA is suspected, despite the absence of evidence. In this study, we described the characteristics and management of PE-related SCA in a large regional registry.

Patients and methods In this prospective population-based study, we included all patients admitted at hospital alive after out-of-hospital SCA, in Paris and suburbs, France (6.6 million inhabitants), from May 2011 to September 2015. Regarding PE, we collected risk factors, clinical decision rules (Wells rule and Geneva score) and diagnostic strategy.

Results Of 2926 patients hospitalized after SCA, 82 cases were diagnosed as PE-related SCA (2.8%, 95% CI 2.2–3.4). Independent factors associated with SCA due to PE were non-shockable initial rhythm (OR 12.4, 95% CI 4.9–31.0, P < 0.001), past history of thromboembolism (OR 10.4, 95% CI 5.6–19.4, P < 0.001), absence of known heart disease (OR 3.8, 95% CI 2.0–7.3, P < 0.001) and female sex (OR 1.9, 95% CI 1.2–3.0, P = 0.008). Considering non-shockable initial rhythm and previous thromboembolism as major predictors of PE, combination of those factors had a specificity for detection of PE-related SCA of 98% and a sensitivity of 23%, with a positive predictive value of 31% and a negative predictive value of 98% (Fig. 1).

Systemic thrombolysis was performed in 47 patients (57%). After adjustment, PE was associated with survival at discharge (OR 2.4, 95% CI 1.2–4.7, P = 0.001), compared with non-PE SCA. Finally, among patients hospitalized for PE-related SCA, only thrombolysis (OR 12.5, 95% CI 1.8–89.1, P = 0.01) and delay from CPR to ROSC < 20 min (OR 6.8, 95% CI 1.3–35.2, P = 0.02) were independently associated with survival to hospital discharge.

Conclusion In this population-based study, PE was not an usual cause of SCA, and was associated with better survival, challenging the traditional view. Thrombolysis was associated with an increased survival in this population, reinforcing current guidelines.

Competing interests None.

O2 Eligibility for and feasibility of donation after circulatory death Maastricht III (DCD MIII) process in post-anoxic patients: a retrospective analysis

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Introduction Donation after circulatory death corresponds to the category III of the Maastricht classification (DCDM III) and may provide mostly kidney and liver transplants with good long-term function. Patients suffering from irreversible brain damages after cardiac arrest are commonly considered candidates for DCDMIII, but little is known regarding the proportion of these patients who could be eligible for this procedure. Using a cohort of post-cardiac arrest patients, our aim was to assess the rate of contra-indications for DCDMIII and to measure the delay between withdrawal of life sustaining treatments (LSTW) and the appearance of low values for common physiological parameters during the agonal phase, which may compromise the process by altering graft function.

Patients and methods Using the Cochin registry (Paris, France), we conducted a retrospective single-centre study from January 2007 to December 2014. We included all patients who died in ICU after LSTW decision because of post-anoxic brain damages. For each patient, we collected exclusion criteria for DCDMIII and the length of time between LSTW implementation and death. We also collected hemodynamic and respiratory parameters during the agonal phase.

Results We included 404 patients in the study, of whom 275 (68%) had at least one exclusion criterion for a DCDMIII process, mostly because of age >65 (190 patients). Other exclusion criteria were: multiple organ failure (n = 88), neoplastic diseases (n = 55, including 46 solid tumours), brain-dead state that occurred after LSTW decision (n = 18), unknown cause of the initial cardiac arrest (n = 13), chronic viral diseases (n = 13), uncontrolled sepsis (n = 4), occurrence of a new refractory cardiac arrest (n = 2), and judicial problems (n = 3). The 130 potentially eligible patients for DCDMII included 94 men (72%) with a mean age of 51 years (±7.7). At time of death after LSTW, the mean length of stay in ICU was 11.6 days (±6). The most common aetiology of cardiac arrest was acute myocardial ischemia (n = 59, 45%). LSTW consisted in terminal weaning of mechanical ventilation in 71 patients (55%), extubation in 12 patients (9%) and infusion of vasopressors was stopped in 3 patients (2%). The average duration of the agonal phase (time between LSTW implementation and death) was 746 min (min) (±162) and this delay was >180 min in 92 patients (71%). After LSTW implementation, an oxygen transcutaneous saturation (SpO2) <70% occurred in 637 min (±545), a mean arterial pressure (MAP) <60 mmHg in 723 min (±586) and a systolic arterial pressure (SAP) <50 mmHg in 733 min (±596). The delay between SpO2 <70% and death was 154 min (±262), and this delay was 59 min (±160) after MAP < 60 mmHg and 23 min (±134) after SAP < 50 mmHg.

Conclusion In this large cohort of brain damaged patients with LSTW decision, we observed that a high proportion of patients would not have been eligible for a DCDMIII process. Even in those without contra-indication, the delay between LSTW implementation and the final circulatory arrest was not compatible with French national guidelines. Low values for arterial pressure and oxygenation persisted during a substantial part of time before final circulatory arrest. This information may help in refining the management of the DCDMIII process in this population.

Competing interests None.

O3 Evaluation of the prognostic value of the bispectral index (BIS) and suppression ratio (RS) among patients admitted to the ICU for cardiac arrest

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Introduction Predicting the neurological outcome of patients admitted to the ICU after a cardiac arrest successfully resuscitated remains difficult [1]. The bispectral index (BIS) allows for the rapid and standardized assessment of the cortical function based upon electroencephalogram analysis whereas the ratio of suppression (RS) is indicative of the absence of electrical activity of the brain. We aimed to evaluate the prognostic value of the BIS and the RS for predicting neurologic outcome after cardiac arrest.

Patients and methods This was a prospective, single center, observational study performed in a large regional University hospital. Adult patients admitted to the ICU for cardiac arrest between March 2012 and October 2014 were included in the study. The exclusion criteria was pregnancy. The BIS and the RS were collected as soon as possible after ICU admission. The patients were not included in the analysis if they died within 24 h, if they had a low signal quality [defined as high EMG artefacts (≥30 dB)], or if the monitoring of BIS started 24 h after ICU admission. The neurological outcome of the patients was based upon the cerebral perfomans category (CPC) calculated at 3 months. CPC score of 1 or 2 indicated good outcome, whereas CPC score of 3–5 indicated poor outcome.

Results During the study period 148 patients were admitted to our ICU for a cardiac arrest. The BIS and RS were monitored in 103 patients.17 patients were excluded (early death ≤24 h; low quality of signal; BIS and RS performed ≥24 h after ICU admission). Thus, 86 patients were enrolled in this study. The mean age was 57.6 ± 16.8 years, 61 patients (70.9%) were male, the cardiac arrest was out-of-hospital in 63 patients (73.3%), hypoxia was the main cause of cardiac arrest (43%), 60 patients (70.6%) were treated with therapeutic hypothermia. At 3 month of follow-up, a total of 50 patients (58.1%) had died and 55 patients (63.9%) were classified as having a poor outcome. The mean duration from the return of spontaneous circulation (ROSC) to the BIS and RS measurements was 5.7 ± 3.0 h. The BIS values were significantly lower in patient with poor outcome compared with patients with a good outcome (5.9 ± 1.1 vs 37.1 ± 18.0, p < 0.0001). The RS values were significantly higher in patient with poor outcome group compared to those with good neurological outcome (85.9 ± 26.3 vs 18.4 ± 31.3, p < 0.0001). The BIS predicted poor outcome with a likelihood ratio of 23.8 and an area under the curve (AUC) of 0.918 [95% CI (0.839–0.966)]. The optimal sensitivity [78.4%, 95% CI (67.3–89.5)] and specificity [96.5%, 95% CI (89.8–100)] for neurological outcome prediction was obtained using a cut-off value of BIS < 5. The RS predicted poor outcome with a likelihood ratio of 23.8 and an AUC of 0.936 [95% CI (0.862–0.977)]. The optimal sensitivity [78.0%, 95% CI (66.9–90.0)] and specificity [96.8%, 95% CI (90.8–100)] for neurological outcome prediction was obtained using a cut-off value of RS > 84. In multivariable logistic regression model, BIS or RS predicted poor outcome with an odds ratio of 65.0 [95% CI (6.1–689.2), p = 0.0005].

Discussion The results of this study using the EEG derived parameters BIS and RS confirm previous findings showing that they are linked to the neurological outcome of patients admitted to the ICU after a cardiac arrest [2]. In particular, a BIS ≤ 5 and/or a RS > 84, measured at ICU admission, were both strongly associated with a poor neurological outcome at 3-months.

Conclusion BIS values may be used to predict long term neurological outcome of patients following cardiopulmonary resuscitation. The ability to accurately predict early non-recovery after cardiac arrest could facilitate discussions with families and limit use of ICU resources in futile cases. BIS and RS values measured at ICU admission might be considered as additional prognostic tools available for the intensivist.

Competing interests None.

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Nutrition during targeted temperature management after cardiac arrest: observational study of neurological and infectious outcomes

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Introduction Cardiac arrest represents one of the greatest medical challenges because of its terrible mortality, morbidity and cost. International guidelines for cardiopulmonary resuscitation led to survival and prognosis improvement, based on the Chain of Survival and the Advanced Life Support, including Targeted Temperature Management (TTM) [1]. Although routinely used, guidelines do not provide detailed management of patients with TTM, especially how to adapt associated therapies such as sedation or nutrition. Guidelines encourage early nutrition for intensive care patients, within 24–48 h after admission [2]. Nevertheless, after cardiac arrest, early nutrition (EN) is disputed. Common post-cardiac arrest syndrome with circulatory failure, frequent diarrheic collapse; and supposed lower digestive tolerance in hypothermia do not encourage this practice. This study first aims to determine if EN is associated with better neurological outcome for patient under TTM. Secondly, we evaluate nutritional tolerance in hypothermia.

Patients and methods We retrospectively included patients under TTM after cardiac arrest in a single mixed intensive care unit from January 2008 to December 2014. Patients fed within 48 h after admission (EN; enteral or parenteral) were compared to those fed after rewarming (DN; after rewarming) concerning neurological and infectious outcomes. Enteral nutrition was initiated at temperature less than 36 °C as compared as ≥36 °C feeding.

Results Among 203 patients under TTM at 33 °C after cardiac arrest, 36 °C feeding. ≥36 °C and <36 °C were recorded for enteral nutrition tolerance comparison between maximal caloric objective defined at 20 kCal/kg/day until day 7, with neurological and infectious outcomes. Enteral nutrition was initiated 48 h or not fed [delayed nutrition (DN); after rewarming] concerning admission (EN; enteral or parenteral) were compared to those fed after cardiac arrest: observational study of neurological and infectious outcomes. Enteral nutrition was initiated 48 h or not fed [delayed nutrition (DN); after rewarming] concerning admission (EN; enteral or parenteral) were compared to those fed after rewarming (DN; after rewarming) concerning admission (EN; enteral or parenteral) were compared to those fed after rewarming (DN; after rewarming). Ninety percent of patients were fed in 33% of the children. Overall survival at 1 month was 8.3% (75/900) (respectively, 27.2% at 36 °C vs 27.3%, p = 0.99; 34 vs 33%, p = 0.87). Conclusion EN is associated with better neurological outcome during targeted temperature management. EN is not associated with more adverse event (infectious and poor nutrition tolerance) when instituted at temperature less than 36 °C.

Competing interests None.

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Description of the epidemiological characteristics and outcome of cardiac arrest in children: a French study

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Introduction Cardiac arrest in children has a very poor prognosis. Knowledge of its epidemiological characteristics is necessary to improve the patient care and survival. The French National Cardiac Arrest registry (RéAC), created in 2009, combines all epidemiological data on out-of-hospital cardiac arrests (OHCA). The objective of this study was to describe the epidemiological characteristics and outcomes of OHCA in children under 18 years old.

Materials and methods All patients under 18 years old victims of out-of-hospital cardiac arrest and registered in the French National Cardiac Arrest registry (RéAC), created in 2009, combines all epidemiological data on out-of-hospital cardiac arrests (OHCA). The objective of this study was to describe the epidemiological characteristics and outcomes of OHCA in children under 18 years old.

Results Out of 42,960 registered cardiac arrest, 900 (2%) involved children under 18 years of age. Out of the 900 patients enrolled, 393 (44%) were less than 1 year old. The OHCA occurred mainly at home (67%). The percentage of traumatic cardiac arrest increased with age, reaching up to 49% in adolescents. Respiratory failure was the leading cause of cardiac arrest in toddlers and children (respectively 40 and 31%). Adolescents were more likely to have an initial shockable rhythm (8%) than other groups (p < 10-3). The intraosseous access was used in 33% of the children. Overall survival at 1 month was 83.3% (75/900) and 66.7% (50/75) of these patients had a favorable neurological prognosis. Outcomes description per age is described in Table 1.

Conclusion There were significant differences between the patient’s groups regarding the location, type of cardiac arrest, initial rate, and survival. An age group approach could be considered to improve care strategy and survival of cardiac arrest victims.

Competing interests None.

Fig. 2 Probability of acquiring VAP according to early or delayed nutrition group

Gray test p = 0.17

- EARLY NUTRITION
- DELAYED NUTRITION

CUMULATIVE INCIDENCE OF VAP

TIME TO VAP (DAYS)
**Table 1 Outcome description of patients**

| Variables | <1 year (n = 393) | 1–4 years (n = 146) | 5–12 years (n = 142) | 13–17 years p (n = 219) |
|-----------|------------------|---------------------|---------------------|-------------------------|
| ROSC | 71 (18) | 43 (30) | 41 (29) | 81 (37) | <10−3 |
| Survival at hospital admission | 68 (17) | 52 (36) | 43 (30) | 85 (39) | <10−3 |
| Survival at day 30 | 13 (3) | 20 (14) | 13 (9) | 29 (13) | <10−3 |
| CPC 1 or 2 at day 30 | 12 (3) | 13 (9) | 3 (2) | 22 (10) | <10−3 |

ROSC return of spontaneous circulation, CPC cerebral performance category

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**O6** Contrast-associated acute kidney injury (AKI) in the intensive care unit (ICU): systematic review and meta-analysis

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Introduction
Avoiding the use of iodinated contrast media (CM) is frequent, fearing it may contribute to AKI. The aim of this systematic review and meta-analysis was to quantify the risk of AKI attributable to CM in ICU patients.

Materials and methods
A systematic review until December 31st 2015, through 5 databases, searched for studies evaluating intravenous administration of iodinated CM. Only controlled studies evaluating AKI following CM exposure in ICU patients matched to unexposed patients were included in the meta-analysis. Meta-analysis was performed on patient-level data using a hierarchical Bayesian nested mixed effects multiple logistic regression model. Bayesian methodology allows evaluating how evidence-based physicians would assess the AKI risk attributable to CM according to both their a priori belief and the presentation of the studies identified in the systematic review. Two meta-analyses were performed with different a priori hypotheses. An objective meta-analysis modeled a neutral state of a priori belief (Odds Ratio [OR] of 1 with impartial distribution) yielding a posteriori OR distribution representative of data collected in controlled studies. A subjective meta-analysis modeled the common belief that CM increases the AKI risk, using an a priori OR of 1.37 based on uncontrolled studies holding clinical community consensus. We determined the minimum a priori relative effective sample size (RESS, representing the a priori strength of belief) needed to observe a significant a posteriori OR distribution: how much physicians have to be convinced a priori that CM increases AKI risk to maintain this belief after being confronted with the studies data.

Results
Among 5696 references, 10 compared ICU patients receiving CM with an unexposed group and 4 performed risk adjustments for baseline AKI risk. Three studies used patient matching: overall, 280 CM patients were matched with 280 control patients. The resulting a posteriori OR did not reach statistical significance: with no prior assumption, there is no evidence that CM increases the risk of AKI in the ICU.

Using an a priori OR of 1.37 (subjective meta-analysis), the a posteriori distribution of the OR did not reach statistical significance except when modeling a very high a priori belief that CM causes AKI (minimum a priori RESS 4.8-folds higher than the RESS of the objective meta-analysis and 70-folds higher than a neutral objective a priori hypothesis).

Conclusion
This systematic review and meta-analysis did not enlighten a risk of AKI attributable to iodinated CM in ICU patients.

Competing interests
None.

O7 Iohexol clearance for exploring the link between glomerular filtration rate and acute kidney injury in patients with acute circulatory failure

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Introduction
Acute kidney injury (AKI) is associated with significant morbidity and mortality, particularly in intensive care unit (ICU) patients. In stable patients, glomerular filtration rate (GFR), the best overall index of kidney function, can be estimated measuring the plasma clearance of an exogenous tracer such as iohexol [1]. There is no reliable method to assess GFR in unstable patients, as classical methods such as MDRD (modifications of diet in renal disease) and urinary clearance calculations have shown their limits [2]. The aim of this study was to determine GFR using iohexol plasma clearance (Clix) at the initial phase of acute circulatory failure and to evaluate its association with subsequent development of AKI.

Patients and methods
Multicentric study in 3 French ICUs. Patients suffering from acute circulatory failure were included within 12 h of ICU admission and administered intravenously a non-toxic dose of iohexol (5 mL; 300 mg/mL) followed by collection of 9 blood samples for iohexol plasma concentration determination (5 and 30 min, 1, 3, 6, 9, 12, 18 and 24 h). Iohexol concentrations were determined using high performance liquid phase chromatography and a three-compartment population pharmacokinetic model was implemented to calculate individual iohexol clearances.

Results
100 patients were included. Median age was 65 years (Q1: 55; Q3: 77), baseline MDRD 93 mL/min (73; 116) and SAPS II 59 (45; 75). Most patients were admitted for septic shock. We could calculate Clix in 85 patients (85%). Failures to calculate Clix included iohexol contrast media injection outside of the study and early renal replacement therapy. Median Clix was 38 mL/min (19–58). 76 patients (76%) developed AKI according to the kidney disease, improving global outcome classification (KDIGO): 15 KDIGO 1, 30 KDIGO 2, 31 KDIGO 3. In 59 patients out of 92 for whom enough serum creatinine dosages were available (64%), serum creatinine decreased in the first 24 h of ICU stay, including 44 patients among the the 76 developing AKI (58%). Clix was inversely related to the severity of AKI: median Clix for KDIGO 0 patients was 68 mL/min (44; 77), 40 mL/min (30; 58) for KDIGO 1 patients, 36 mL/min (24; 52) for KDIGO 2 patients and 16 mL/min (9; 22) for KDIGO 3 patients. In 40 out of 82 patients (49%) the difference between MDRD (calculated from serum creatinine at the time of inclusion) and Clix exceeded 20 mL/min, and in most cases (90%) MDRD overestimated estimated GFR (eGFR). For
AKI patients, eGFR according to MDRD was >60 mL/min in 18 patients. According to MDRD, 8 patients had glomerular hyperfiltration defined by an eGFR > 130 mL/min. Only 4 patients had hyperfiltration according to Cioxx. For the 2 patients with eGFR > 130 mL/min with both methods, one suffered denutrition; his Cioxx was 138 mL/min, while MDRD was 281 mL/min. For the 6 patients with hyperfiltration according to MDRD and not Cioxx, 4 had denutrition, 1 had a very low baseline serum creatinine (18 μmol/L), and one suffered morbid obesity. Four patients with eGFR > 130 mL/min according to MDRD had a maximum KDIGO score of 2.

**Discussion** Our study confirms that variations of serum creatinine are not a good marker of GFR. We hypothesize this to be related to the large amount of fluid infusion at the acute phase following ICU admission and the influence of nutritional factors. The MDRD formula tended to overestimate eGFR. Cioxx may enable to overcome the limits of the MDRD formula at the acute phase of critical illness, as it seems not to be influenced by nutritional factors or fluid infusion, unlike creatinine variations.

**Conclusion** The close link between Cioxx and AKI is very encouraging for the development of this method of eGFR assessment in critically ill patients. Cioxx could be used for early determination of AKI risk and for drug dosage adaptation, as it is a better reflection of instantaneous GFR than MDRD.

**Competing interests** None.

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**O8**

**Effect of renal replacement therapy strategies in septic-shock patients with severe acute kidney injury: a post hoc analysis of a randomized controlled trial**

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**Introduction** Acute kidney injury is particularly common in septic-shock patients and is associated with high mortality. The putative effect of renal replacement therapy (RRT) on the prognosis of such patients is highly debated; some advocating that outcome might improve owing to modulation of inflammation. We aimed to compare outcomes of septic-shock patients with severe acute kidney injury (stage 3 of KDIGO classification) treated with an early RRT strategy (all patients immediately received RRT) with those treated with a delayed RRT strategy (patients received late RRT or no RRT at all).

**Patients and methods** We did a post hoc subgroup analysis in a subset of septic-shock patients with severe acute kidney injury (stage 3 of KDIGO classification) from a multicenter randomized controlled trial. In the trial, patients from 31 intensive care were randomly assigned (1:1) to either an early or a delayed RRT initiation strategy. With the early strategy, RRT was initiated within 6 h after inclusion criteria were met. With the delayed strategy, RRT was started if at least one of the following criteria was met: severe hyperkalemia, metabolic acidosis, pulmonary edema, serum urea concentration greater than 40 mmol/l, or oliguria for more than 72 h after randomization. The primary outcome was overall survival at day 60.

**Results** Of the 413 septic-shock patients (on a total of 620 patients), 209 were managed with early strategy and 204 with delayed strategy. The Kaplan–Meier estimates of mortality at day 60 did not differ significantly between the early and delayed strategies; 101 deaths occurred among 209 patients in the early-strategy group (48.5%; 95% confidence interval [CI] 41.3–54.9), and 99 deaths occurred among 204 patients in the delayed-strategy group (48.5%, 95% CI 41.2–55.0; P = 0.97) (Fig. 3). A total of 97 patients (47.5%) in the delayed-strategy group did not receive renal-replacement therapy. The number of days RRT-free days was significantly higher in the delayed strategy group (21 [5–29] vs 17 [2–25], p < 0.001). Median length of stay in hospital did not differ significantly between groups (20 [8–39] vs 19 [7–40] days, p = 0.9).

**Conclusion** The timing of RRT in septic-shock patients with severe acute kidney injury did not significantly influence mortality. However, a conservative strategy avoided many unnecessary RRT sessions.

**Competing interests** None.

**O9**

**Dose response multicentre investigation on fluid assessment (DOREMIFA) in critically ill patients: the French cohort**

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Annals of Intensive Care 2017, 7(Suppl 1):O9

Introduction There is growing evidence that fluid accumulation beyond the correction of hypovolemia is associated with increased morbidity, mortality and a longer hospital stay. We recently published a prospective cohort observational study “The Dose Response Multicentre Investigation on Fluid Assessment (DoReMIFA) in critically ill patients” aimed to investigate the impact of fluid balance and fluid accumulation on mortality for both AKI and (Non) N-AKI patients as for those who receive renal replacement therapy (AKI-RRT).

Aim of the present is to assess the fluid administration of the French subgroup of patients.

Patients and methods We analysed 209 (12.05%) of the 1734 enrolled patients, from the 2 French ICUs. Fluid overload (FO) was defined as the ratio between cumulative fluid balance and the initial body weight, in percentage. Maximum fluid overload (MFO) referred to the peak value of FO during the entire ICU stay. TMFO represented the number of days between ICU admission and day of MFO. Velocity of fluid accumulation was defined by Fluid overload slope (FOSL) as the MFO/TMFO ratio. A boxplots for the three groups (N-AKI, AKI and AKI-RRT) illustrated the MFO for both survivors and non-survivors during the ICU stay. A Kaplan-Meier analysis was performed to evaluate the time to death for the three groups (N-AKI, AKI and AKI-RRT). The time to death was evaluated by a Cox proportional hazard regression analysis.

Results 53% of patients had AKI (38% stage 1, 20% Stage 2, 42% Stage3). The Kaplan-Meier analysis including the first 28 days of ICU, highlighted a significant survival benefit for patients without AKI, in particular for longer ICU stay. The AKI and AKI-RRT group had, conversely form the entire study population, similar survival rates. In all cohorts as in N-AKI, AKI and AKI-RRT non-survivors had a higher MFO than survivors. Again, the AKI-RRT and AKI groups had similar levels of MFO, with a lower over-hydration for the RRT group. Cox regression analysis of the velocity of fluid accumulation showed that for every increase of one unit of the FOSL, the hazard of death increased significantly by a factor of 1.44. The hazard ratio decreased to 1.41 when adjusting for SAPS II score.

Discussion The fluid assessment in critically ill patients enrolled on the French ICUs confirm the findings of the Doremifa study. Fluid overload is strongly correlated with mortality at any degree. A lower degree of fluid overload for the AKI-RRT group, when compared with the AKI population, and a similar survival curve between the 2 groups, may suggest that CRRT has a protective effect. More analysis are needed to confirm this hypothesis, an early initiation of the treatment have to be also investigated.

Conclusion The velocity of fluid accumulation, as in the findings of the main study, contribute to worse patients outcome.

Competing interests Fresenius medical care: fees for travelling and hotel.

O10 Doppler-based renal resistive index in assessing renal dysfunction reversibility in ICU patients: results of a multicenter cohort study

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Introduction Doppler-based renal resistive index (RI) measurement may hold promise in differentiating transient from persistent AKI in selected critically ill patients. Although several studies have suggested adequate performance in predicting short-term reversibility of AKI, most of these studies were performed in limited patient samples [1]. Additionally, a recent study has identified discrepant results regarding its diagnostic performance [2] suggesting confirmatory studies to be required.

The main objective of this study was to assess diagnostic performance of RI in predicting persistent AKI in critically ill patients. Secondary objectives were to assess diagnostic performance of semi-quantitative assessment of renal perfusion (SQP) using color-Doppler in predicting persistent AKI and performance of both tests in predicting needs for renal replacement therapy (RRT).

Patients and methods Prospective multicenter study performed in eight ICUs from December 2013 to April 2016. This study was declared to Clinicaltrial.gov.

Adult patients requiring mechanical ventilation were included in this study. Patients with mild to severe chronic kidney diseases, arrhythmia, or obstructive renal dysfunction were excluded from this study. Patients with hospital stay shorter than 72 h and in whom renal reversibility could not be assessed were secondarily excluded.

Acute kidney injury (AKI) was defined according to the KDIGO definition. Transient AKI was defined as AKI with recovery within the first 3 days following inclusion.

Intra-renal RI was calculated as (peak systolic velocity − end-diastolic velocity)/peak systolic velocity.

SQP was assessed using a scale ranging from 0 (absence of renal perfusion) to 3 (renal vessels identifiable until the arcuate arteries in the entire field of view).

Results are reported as number (%), median (IQR) and area under curve (95% CI).

Results Overall, 371 patients were included. Median age was 76 years (66–89) and 236 patients were of male gender (63.6%). Most of the patients were admitted with medical conditions (n = 253; 68.2%) and 162 patients (43.7%) had sepsis at ICU admission. Median LOD score was of 8 (5–11) at study inclusion and 198 patients required vasopressors (53.4%).

Of the included patients, 253 (68.2%) had an AKI at study inclusion, including 158 patients (42.6%), 35 (9.4%) and 60 (16.2%) with AKI stage 1, 2 and 3 respectively. Doppler-based RI was obtained in 365 patients (98.4%), semi-quantitative assessment of renal perfusion in 367 (98.9%), Patients with AKI had a higher RI at ICU admission [0.70 (0.62–0.77) vs. 0.65 (0.59–0.70); P = 0.0001] and a lower SQP [2 (1–3) vs. 2 (2–3); P = 0.0003].

Twenty patients were discharged before day 3 leaving 351 patients in the final analysis, including 118 (33.6%), 97 (27.6%) and 136 patients (38.7%) no AKI, transient AKI or persistent AKI respectively. Resistive index at inclusion was of respectively 0.65 (0.59–0.70), 0.69 (0.62–0.77) and 0.71 (0.62–0.77) in patients without AKI, with transient AKI and with persistent AKI (P = 0.0005). Resistive index failed to demonstrate any interest in predicting persistent AKI (Area under ROC curve: 0.58; 95% CI 0.52–0.64).

Semi-quantitative assessment of renal perfusion was of respectively 2 (2–3), 2 (2–3), and 2 (1–3) in patients without AKI, with transient AKI and with persistent AKI (P = 0.002). Semi-quantitative assessment of renal perfusion failed to demonstrate any interest in predicting persistent AKI (Area under ROC curve: 0.59; 95% CI 0.52–0.65).

Overall, 46 patients (12.4%) required RRT during ICU stay. In these patients, RI and SQP were of respectively 0.75 (0.65–0.77) [vs. 0.67 (0.60–0.74) in non-RRT patients; P = 0.0003] and 2 (1–2) [vs. 2 (2–3) in non-RRT patients; P = 0.0003]. Both test displayed a poor performance in predicting subsequent renal replacement therapy at study inclusion.
A poor functional outcome was reported for 55/90 (61%) patients at 90 days (5 patients were lost to follow-up). The following factors were independently associated with a poor outcome at 90 days: older age (per 1-year increment, adjusted odds ratio (aOR): 1.04, 95% CI 1.0–1.08, \( p = 0.035 \)), cerebrospinal fluid (CSF) protein level >1.9 g/L (aOR: 8.85, 95% CI 2.49–39.64, \( p = 0.002 \)), hydrocephalus on MRI (aOR: 8.6, 95% CI 2.05–46.5, \( p = 0.006 \)). By contrast, the use of adjunctive steroids had a protective effect (aOR: 0.15, 95% CI 0.02–0.87, \( p = 0.045 \)). The Kaplan–Meier estimated 1-year mortality was 51% (0.39–0.61). The following factors were independently associated with mortality: CSF protein levels >1.9 g/L [adjusted relative risk (aRR): 2.47, 95% CI 1.17–5.25, \( p = 0.018 \)], hydrocephalus on MRI (aOR: 3.45, 95% CI 1.50–7.91, \( p = 0.003 \)), brain infarction on MRI (aRR: 2.40, 95% CI 0.99–5.81, \( p = 0.051 \)). The use of adjunctive steroids had a protective effect on 1-year mortality (aRR = 0.16, 95% CI 0.05–0.45, \( p = 0.0006 \)).

**Conclusion** Despite antituberculous therapy and supportive care, severe forms of TBM are characterized by a poor outcome in more than 50% of cases. Elevated CSF protein levels, hydrocephalus and brain infarction on MRI at ICU admission represent major indicators of poor outcome. Our data suggest that use of adjunctive steroids is associated with reduced disability and mortality, irrespective of immune status and severity of illness. We conclude that adjunctive steroids may benefit all patients with a suspicion of TBM admitted to the ICU.

**Competing interests** None.
AUC at 0.86. RHDE during deep exhalation predicts a peak expiratory flow <270 l/min with AUC at 0.78.

**Conclusion** Diaphragmatic echography may predict respiratory insufficiency in neurosurgical disorders.

**Competing interests**

None.

### O13 Brain injury during veno-arterial extracorporeal membrane oxygenation

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**Introduction** Veno-arterial extracorporeal membrane oxygenation (VA-ECMO) is used to provide cardiac support in patients suffering refractory cardiogenic shock. Its use is increasing and associated with neurological complications, mainly cerebrovascular. The frequency of those events and their impact on patients are not well described. We therefore study the epidemiology, risk factors and impact of cerebral complications occurring in VA-ECMO patients.

**Patients and methods** Observational study conducted in a tertiary referral center (2006–2014) on patients developing a neurological complication (ischemic stroke or intracranial bleeding) while on VA-ECMO versus those who did not.

| Characteristic | No brain damage | Ischemic stroke (n = 47) | Cerebral bleeding (n = 25) |
|---------------|-----------------|-------------------------|---------------------------|
| Age (years)   | 50 ± 15         | 50±16                   | 48 ± 18                   |
| Male sex (%   | 568 (71)        | 32 (68)                 | 12 (48)                   |
| SAPS II       | 72 [54–85]      | 70 [61–80]              | 74 [57–90]                |
| SOFA score    | 5 [3–14]        | 5 [3–8]                 | 3 [3–7]                   |
| Biologic values (J1 ECMO) | | | |
| Lactates (mmol/L) | 6 [2.7–11.1]    | 5 [2.9–10.1]            | 4 [4.0–10.0]              |
| PT (s)        | 45 [30–62]      | 54 [34.65]              | 47 [27.5–54]             |
| ACT           | 1.6 [1.2–2.3]   | 1.4 [1.1–1.8]           | 1.8 [1.4–3]              |
| Fibrinogen (g/L) | 3.0 [2.0–5.0]   | 3.0 [2.4–5.0]           | 4.0 [1.7–4.0]            |
| Platelets (×10¹²/L) | 155 [96–215]   | 163 [100–246]           | 86 [56.5–164]            |
| Type of ECMO  |                 |                         |                           |
| Peripheral    | 698 (87)        | 35 (74)                 | 20 (80)                   |
| Switch to central ECMO | 38 (5)  | 2 (4)                   | 4 (16)                    |
| Central       | 107 (13)        | 12 (26)                 | 5 (25)                    |
| Intra-aortic balloon pump | 265 (33) | 16 (34)                 | 5 (25)                    |

Results are expressed as mean ± SD, number (%) or median [27th–75th percentile interquartile range]

**Results** The main clinical characteristics of the patients are reported in Table 2. Among 873 consecutive patients who had received VA-ECMO, 72 developed cerebral complications on ECMO: ischemic stroke in 47 (5%) and cerebral bleeding in 25 (3%), occurring after a median (IQR) of 11 (7–21.5) and 6 (5–11) days of ECMO support, respectively. No specific risk factor of ischemic stroke was found in univariable analysis except body mass index >26 (OR 2.16, 95% CI 1.15–4.05). Hematological failure (defined as platelets <50 000/mL at ECMO initiation (OR CI 3.64, 95% 1.30–10.21) and platelets <20,000/mL during ICU stay (OR 3.01, 95% CI 1.22–7.40) were significantly associated with cerebral bleeding in univariable analysis. Age, comorbidities, renal replacement therapy, and intra-aortic balloon pump use were not associated with neurological complications. Twenty-three (49%) patients with ischemic stroke and 21 (84%) with intracranial bleeding died versus 385 (48%) of patients without brain injury.

**Conclusion** Neurological events occurred frequently in patients on VA-ECMO. Ischemic stroke is the most frequent, occurs late during ECMO support and does not seem to be associated with higher mortality than patients without brain injury. Cerebral bleeding occurs early and is associated with high mortality rate. Low platelets count at ECMO initiation and during ECMO support are associated with cerebral bleeding.

**Competing interests**

None.

### O14 Does heart rate variability predict clinical outcome of patients with subarachnoid hemorrhage in the neurointensive care unit

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**Introduction** Patients with subarachnoid hemorrhage (SAH), admitted to the neurointensive care unit (NICU) are exposed to complications including rebleeding, vasospasm, hydrocephalus, pain and sepsis. The autonomic nervous system (ANS) is a warning system which can be assessed noninvasively by heart rate variability (HRV). Many reports have shown a relationship between HRV and outcome in myocardial infarction, stroke and renal insufficiency. To our knowledge very few papers have addressed this issue in the NICU. The aim of this study was to check whether HRV could predict outcome in patients with SAH admitted to the NICU.

**Patients and methods** Following Institutional Review Board approval, patients with SAH, admitted to the NICU were included in this prospective monocentric study. Those with persistent arrhythmia, cardiac pacing or younger than 18 years were excluded. All subjects were assessed everyday starting from their arrival at NICU, i.e. day 2 after SAH, to day 7. HRV was measured between 2 and 4 p.m. during 10-min. HRV was achieved by connecting a computer to the electrocardiogram (ECG) monitor. Online power spectrum was calculated from the ECG R–R interval using the maximum entropy method (MemCalc™, Suwa Trust, Japan). Low (LF: 0.04–0.15 Hz) and high frequency (HF: 0.15–0.4 Hz) spectra were associated with sympathetic (Σ) and vagal activities respectively. Entropy and coefficient of variation of RR intervals (CVVR) were also measured. Concomitantly, we noted demographic, hemodynamic, respiratory and comorbidty data. The severity of SAH was classified using the five points World Federation of Neurological Surgeons (WFNS) score (were 1 = less severe and 5 = very severe) and the four point Fisher scale (where 1 = less severe and 4 = very severe). Outcome at discharge was assessed by the modified Rankin score, a poor outcome was defined by either death or Rankin score of 4–5 which means severe disability and inability to walk without assistance Outcome was defined as good or poor based on the absence or presence of Rankin score of 4–5 and death. Univariate and multiple logistic regression models were applied on these data to determine the predictive (s) factor (s) of poor outcome.
Results
The inclusion criteria were fulfilled by 125 patients, but complete data was only available in 53 of them. Among them, five died before discharge, 10 were discharged with a Rankin score of 4 or 5. The most significant modifications of our study parameters were observed on day 2 after SAH. These observations are summarized in Table 3. Consequently, on day 2 after hemorrhage, LF (p = 0.02), VF (p = 0.03), entropy (p < 0.001) and CVVR (p < 0.01) were significantly decreased in patients with poor prognosis. This trend was sustained The multivariate logistic regression model revealed an odds ratio (95% confidence interval) for LF = 0.997 (0.995–1.000); Entropy = 0.919 (0.872–0.968); VF = 0.997 (0.995–1.000) and CVVR 0.586 (0.388–0.885) considered as significant predictors of poor outcome.

Discussion
These results corroborate previous reports showing that a decrease in HRV, particularly the sympathetic pathway, at admission, is associated with poor outcome (1).The mechanisms are not clearly known. Ischemic or hemorrhagic lesions had been observed in the hypothalamus of patients with poor outcome after SAH, which is the predominant modulator of autonomic system (2).

Conclusion
This study suggests that early assessment of HRV in the NICU could predict outcome in patients with SAH. A decrease in HRV at admission of patients was significantly associated with poor outcome. Further studies are required to confirm this finding.

Competing interests
None.

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O15
Electrical muscle stimulation and bicycling combined to early standard rehabilitation versus early standard rehabilitation alone: impact on global muscle strength at ICU discharge—an open-label, single-centre, assessor-blinded randomised trial
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Introduction
Early Standard Rehabilitation (ESR), first passive and then passive/active, is recommended for critically ill patients in whom it reduces the duration of mechanical ventilation (MV), improves functional status, muscle strength and quality of life after hospital discharge. The early addition of leg bicycling on a cyclo-ergometer and of electrical muscle stimulation (EMS) is now part of common practice in the ICU. Whether it can preserve or improve muscle strength and further increase the beneficial effects of ESR is little known.

Patients and methods
Single-centre, randomised study comparing the effects of the combination of early and daily leg bicycling + EMS of the quadriceps + ESR (intervention group) versus ESR alone (usual care group) on the global muscle strength assessed by the MRC score at ICU discharge by a physiotherapist blinded to the randomization group (NCT02185989). All consecutive patients were potentially eligible if they were deemed to need more than 72 h of care in ICU. Main non-inclusion criteria were resuscitated cardiac arrest, presence of pacemaker or implantable defibrillator, acute cerebral disease requiring deep sedation for at least 72 h, known neuromuscular disease, and amputation of a lower limb. Randomization was stratified by sex, MV or not at study entry, and day of admission (Thursday/Friday or other days). The interventions were applied right from Day 1 (within 72 h of admission), 5 days/week. Protocolled ESR consisted of daily multistep program (from 10 passive mobilisations of each joint in comatose patients to passive/active muscle work, transfer to chair, standing and walking, depending on patient’s level of wakefulness/ cooperation). In the intervention group, 30 min passive/active leg bicycling (even in bed-ridden patients) and 54 min EMS of the quadriceps were performed 5 days/week in addition to ESR, according to pre-established programs.

Results
From July 2014 to June 2016, 314 patients were included (as planned per protocol) and 313 were analysable (1 consent withdrawal): 155 in usual care group and 158 in intervention group. Among the whole population, ICU mortality was 18%, SAPSII 46 ± 18, admission SOFA 8 (IQR 6; 12), patients treated with MV 85%. Clinical characteristics at study entry were similar between groups. Primary endpoint: 124 and 121 patients upon the 131 and 125 ICU survivors in usual care and intervention groups respectively, could be assessed for the primary endpoint. The discharge MRC score was 53 (IQR 44; 60) and 51 (44; 58) in usual care and intervention groups respectively (P = 0.86), and was also not different between groups in patients under MV at time of study entry: 52 (IQR: 44; 58) (n = 95) and 49 (43; 57) (n = 89) (P = 0.26).

Secondary endpoints: There was no between-group difference in discharge functional status as assessed by the ICU mobility scale in the whole population (P = 0.54) (P = 0.64 in patients under MV at study), by the change in the Katz index from inclusion to discharge (P = 0.39) (P = 0.47 in patients under MV at study entry), or in the day-28 ventilator-free days [21 days (17; 22) vs 20 days (18; 21); P = 0.34]. The thickness of the rectus femoris muscle, assessed by echography at inclusion and discharge in survivors, showed a lower decline in the intervention group: −1.8 mm (−4.4; −0.2) vs −0.7 mm (−1.5; −0.25); P = 0.009. The impact on delirium occurrence in ICU is still under analysis. Data concerning physical and mental status at 6 months are not fully available yet.

Safety: We observed no serious adverse event related to the studied interventions.

Conclusion
Although safe and resulting in lower decline in muscle thickness as observed on echography (not blinded assessment), the addition of daily leg bicycling and EMS to ESR did not result in higher global muscle strength as assessed by the MRC score (blinded assessment) at ICU discharge in a mixed and heterogeneous population of critically ill patients. Exploratory subgroup analyses are underway and perhaps will help to identify subsets of patients in whom the studied intervention might be beneficial and might deserve further investigations.

Competing interests
None.

O16
Post-intensive care syndrome: a population-based observational study of healthcare use
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Table 3  Univariate analysis of data in the study population

|                      | Good outcome | Poor outcome | p value |
|----------------------|--------------|--------------|---------|
| No of patients (%)   | 38 (72)      | 15 (28)      |         |
| Mean age (years)     | 56.2 ± 11.7  | 59.6 ± 12.1  | 0.004   |
| Female sex           | 23 (60)      | 9 (60)       | 0.86    |
| WFNS IV–V            | 10 (26)      | 7 (47)       | <0.001  |
| Fisher grade 4       | 18 (47)      | 9 (60)       | <0.001  |
| HRV on day 2         |              |              |         |
| LF (ms)              | 519 ± 528    | 171 ± 385    | 0.02    |
| HF (ms)              | 513 ± 863    | 224 ± 505    | 0.24    |
| Entropy              | 50.3 ± 1.20  | 34.3 ± 16.4  | <0.001  |
| LF/HF                | 1.97 ± 1.41  | 1.87 ± 2.27  | 0.85    |
| VLF (ms)             | 1471 ± 2152  | 184 ± 364    | 0.02    |
| CVVR (%)             | 4.18 ± 2.01  | 2.25 ± 2.31  | 0.006   |
Introduction Intensive care unit (ICU) admission is known to lead, among survivors, to numerous and persistent disabilities and impairments after discharge, forming the "post intensive care syndrome" (PICS). However, PICS consequences in term of healthcare use is less studied. Recent data tend to demonstrate that ICU survivors have increased healthcare use after ICU. However, little is known about healthcare use for the most severely ill patients, which are supposed to be at higher risk of PICS, and, to our knowledge, no epidemiological data are available in France.

Patients and methods We conducted a retrospective multicenter study using comprehensive administrative hospital discharge databases of the Centre Val de Loire region, France (2.5 millions of inhabitants). Based on an ICD-10 algorithm, we included all adult patients admitted in an ICU for septic shock or acute respiratory distress syndrome (ARDS) during 2011 and invasively ventilated at least 5 days. Performance of the selection algorithm was validated through review of a subsample of medical charts. Comorbidities were also extracted from ICD-10 coding and reported using a scoring system derived from Charlson Comorbidity Index. Healthcare use and comorbidities were analyzed 2 years before (pre-ICU period) and 2 years after ICU (post-ICU period).

Results 552 patients were selected, of which 249 (45%) died during the hospital stay. Among the 303 survivors, 293 (97%) had complete data required for analysis and none was lost for follow up. Mean ± SD age was 61 ± 14 years, SAPS2 49 ± 17 and median ventilation duration was 10 days (Q1 = 7; Q3 = 20). Regarding chronic comorbidities during the pre-ICU period, cardiac disease was reported for 26% of the patients, respiratory disease for 16%, kidney disease for 13%, and hepatic disease for 12%.

Healthcare resources utilization analysis during the pre-ICU period revealed that 58% of the patients required hospitalization, 54% ambulatory care, 57% emergency admissions and 10% rehabilitation facilities. Twenty-three percent of the patients had no healthcare use. During the post-ICU period, the 2-year mortality rate was 15%. Healthcare resources utilization was significantly increased during the post-ICU period compared to the pre-ICU period for hospitalizations (72%, p < 0.001), ambulatory care (73%, p < 0.001) and rehabilitation facilities (54%, p < 0.001). No patient had no healthcare use. Regarding chronic comorbidities, cardiac, respiratory and renal diseases were significantly more frequent compared to the pre-ICU period (respectively 32, 27, and 21%, p < 0.001 for the three conditions). Time trend analysis of the healthcare use in the post-ICU period revealed that the first 9 months were at high healthcare use (essentially hospitalizations and rehabilitation facilities), and emergency admissions tended to increase at the end of the 2-years follow-up.

Discussion Patients admitted to ICU for acute respiratory distress syndrome and septic shock frequently have a significant healthcare resources utilization during the 2 years before. The 2 years following admission is characterized by a more important healthcare use, together with a significant increase in comorbidities.

Conclusion Our study highlights the epidemiological impact of PICS at the population level in a French region, underpinning observational and interventional research within and beyond the ICU.

Competing interests None.

O17 A pilot study of 6-months evaluation of social, psychological, financial and emotional consequences of an ICU stay in survivors critically ill patients Cécile Rebière1, Elie Azoulay2, Benoit Misset3, Stephane Ruckly4, Jean-François Timst5, Maitë Garrouste-Orgeas1
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Table 4 Functional and personal changes in cases and controls

| Variables                          | Cases n = 37 | Control n = 37 | p value |
|------------------------------------|-------------|----------------|---------|
| Return home within 1 month         | 17 (45.9%)  | 37 (100%)      | <0.01   |
| Need of help at home               | 22 (59.4%)  | 5 (13.5%)      | <0.05   |
| Change in working conditions       | 12 (32.4%)  | 3 (8.1%)       | <0.05   |
| Change in financial conditions     | 10 (27%)    | 7 (18.9%)      | 0.6     |
| Psychologist help                  | 17 (45.9%)  | 5 (13.5%)      | <0.05   |
| Use of medications                 | 9 (24.3%)   | 8 (21.6%)      | 1       |
| Change in marital status           | 2 (5.4%)    | 3 (8.1%)       | 1       |
| Eating disorders                   | 8 (21.6%)   | 5 (13.5%)      | 0.54    |
| Disorder of sexual life            | 12 (32.4%)  | 7 (18.9%)      | 0.3     |

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Table 4 Functional and personal changes in cases and controls

Introduction Consequences of hospitalization in critically ill patients have been recognized for several years with physical, cognitive and psychological consequences published under the denomination of the post intensive care syndrome which will become a new challenge for intensivists. The impact of social, personal and financial consequences has been less reported. The primary objective of this pilot study is to report the social, financial, psychological, physical and emotional consequences in a group of critically ill patients compared to a group of patients never hospitalized in ICU. The second objective is to investigate patient's perceptions to better understand their memories of their hospitalization through a qualitative approach.

Patients and methods We designed a case control study in three ICUs belonging to the Outcomerea research group (July 2014–May 2015). Case patients were adult patients ventilated for more than 48 h. We excluded patients not speaking or understanding French and patients who denied participation. Clinical and demographics characteristics of the cases were extracted from the Outcomerea database. They were interviewed 6-months after ICU discharge. Control patients, matched on age and sex and never hospitalized in ICU, were interviewed face-to-face during an hospital consultation. All patients completed the same questionnaires in a random order, exploring emotional and post traumatic-stress syndrome (Impact of Event Scale-revised, cut-off >22), self-sufficiency in daily activities (activity of daily Living, ADL), quality of life (first question of the SF-36), and questions about their place of living, of working and financial conditions, need of psychological help and marital status. We used a phenomenological approach to report patient’s perceptions.

Results Of the 96 eligible patients, 20 (20.8%) died at 6 months, 39 (40.6%) were excluded and 37 were entered in the analysis and compared to 37 control patients. Characteristics of the case patients were: age (median: 65, range: 47–73), 64% male, SAPS II (51, 37–64), ICU stay (12 days, 8–19), hospital stay (29 days, 22–41). The median IES-R score was significantly higher in cases (14, 8–31) vs control (6, 3–10), p < 0.01. IES-R > 22 was found in 13 (35.1%) cases patients versus 1 (2.7%) of control patients (p < 0.01). Activities of daily living without help were significantly more often performed in control versus cases for bathing (n = 37, 100% vs n = 33, 89%, p = 0.04) and continence (n = 37, 100% vs n = 33, 89% p = 0.04). Perception of the quality of life was not significantly different between cases and controls but increasing quality of life was much important in cases (n = 15, 40.6%) versus control (3, 8.1%), p < 0.01. See Table 4.
Three themes were found in the qualitative analysis: the ICU stay seen as a traumatized period, a period without memories and support from families and friends. Their representations in the verbatims were 13 (35.1%), 13 (35.1%) and 3 (8.1%).

**Conclusion** This pilot study reported substantial neuropsychological and functional alterations related to the ICU stay and emphasized the need for better estimating these modifications in a multicenter study. Addressing these consequences adds to the role of intensivists for elaborating prevention programs and promoting post intensive care syndrome to non ICU practitioners to collaborate together for the best future of ICU patients.

**Competing interests** None.

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**O18**

**To understand or not to understand brain death: impact on grief symptoms in relatives with experienced organ donation request**

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**Annals of Intensive Care 2017, 7(Suppl 1):O18**

**Introduction** In the ICU context, in the case of organ donation, patients’ relatives are at the centre of the decision process: within a limited time frame, they will be told that the patient is brain dead and will be asked to consider organ donation. Qualitative studies have put forward that understanding brain death facilitates decision-making and impacts on the final decision (donation vs non donation). However the impact of understanding brain death on relatives’ grieving process has never been evaluated. In this study, we searched for correlation between semi-quantitative answers to questions related to understanding of brain death and experience of the process in a questionnaire completed by relatives 1 month after the patient’s death and post-traumatic related symptoms (PTSD) and complicated grief.

**Patients and methods** This is an ancillary study of a larger prospective, observational study in 28 ICUs in France that aimed to compare grief symptoms of relatives of donor patients versus relatives of non-donor patients. For each brain dead patient, the relative who served as the surrogate was included at time of organ donation discussion. Relatives were assessed at 3 time points during a telephone interview: at 1 month, to complete a questionnaire regarding their experience in the ICU and description of the organ donation request and procedure, including understanding of brain death; at 3 months to complete the Hospital Anxiety and Depression Scale (HADS) and the Impact of Event Scale-Revised (IES-R) for PTSD symptoms; at 9 months, to complete the IES-R and the Inventory of Complicated Grief (ICG).

**Results** 202 relatives were included in the study. At 1 month after the patient’s death, 79.2% of relatives completed the questionnaire, at 3 months 70.3% completed the HADS and the IES-R and at 9 months 61.4% completed the IES-R and the ICG.

One month after the death, 35% of relatives declared having difficulties in understanding brain death and 32% experienced decision-making as difficult. Results show that experience of the decision making process impacts on relatives’ well-being. At 3 months, compared to relatives who did not find the decision difficult, those who did find it difficult more often presented significant PTSD symptoms (40.54 vs 65.88%, p = 0.016). At 9 months, compared to relatives who understood brain death, those who did not understand brain death had higher global IGC score [23 (12.5–36.5) vs 36 (28–43.75), p = 0.010] and more often presented complicated grief symptoms (46.15 vs 75%, p = 0.026). There was a trend in increased prevalence of PTSD related symptoms with 60% in the group of relatives who did not understand brain death versus 47.2% in the group that did, but this was not significant (p = 0.33).

**Discussion** Results show that difficulty experienced during organ donation discussion and decision impacts on relatives’ well-being in the months that follow the patient’s death. Support to relatives should be proposed in this context. Interestingly, understanding of brain is a key component of relatives’ experience: on top of possibly impacting on the decision itself, it significantly impacts on relatives’ grieving process 9 months after the patient’s death. Promoting better understanding of brain death, proposing clearer explanations, by using various media, may improve both relatives’ understanding and well-being.

**Conclusion** Our study shows that understanding of brain death is a key component of relatives’ experience that significantly impacts on the grieving process. Efforts should be made to improve relatives’ understanding of brain death.

**Competing interests** None.

**O19**

**Impact of ICU end-of-life care on relatives’ grief symptoms**

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**Annals of Intensive Care 2017, 7(Suppl 1):O19**

**Introduction** Relatives of patients who die in the ICU experience a considerable burden of harm such as symptoms of anxiety and depression, posttraumatic stress disorder (PTSD) symptoms and quality-of-life alterations. Improving the quality of dying and death
is recognized as a priority. Nevertheless, specific data are needed to understand what specific aspects of ICU care affect the relatives’ grieving process. This study aims at providing information on potential links between anxiety/depression, PTSD related symptoms, complicated grief and components of ICU end-of-life care in order to specify ICU practices that may affect the risk of developing these symptoms.

**Patients and methods** This is an ancillary study of the CAESAR study—a prospective, observational study in 41 ICUs in France. Eligible patients were adults who died after at least 48 h in the ICU. For each patient, the relative who served as the surrogate was included at time of death. Relatives were assessed 21 days then 3, 6, and 12 months after the death during a telephone interview. At 21 days they completed the CAESAR scale; at 3 months, they completed the Hospital Anxiety and Depression Scale (HADS) and the Impact of Event Scale-Revised (IES-R) and at 6 and 12 months they completed the IES-R and the Inventory of Complicated Grief. In this study, we searched for correlation between semi-quantitative answers to CAESAR questions (and not scores) and outcomes.

**Results** 475 patients and their relatives were included. Response rates were 90.5, 81.3, 59.4 and 45.2%, at day-21, 3, 6 months and at 1 year, respectively. 5 domains are associated with significant increased risk of developing ICU burden (p ≤ 0.05 for each variable).

1. **Quality of care and symptom control.** Perception that pain was not under control and that the patient had difficulties in breathing is associated with increased risk of developing anxiety and depression at 3 months, PTSD related symptoms at 6 and 12 months, complicated grief at 6 months. Dissatisfaction with quality of care is associated with increased risk of developing anxiety and depression at 3 months, PTSD related symptoms at 3 and 6 months, complicated grief at 6 months.

2. **Quality of communication.** Dissatisfaction with communication with either doctors or nurses is associated with increased risk of developing anxiety and depression at 3 months, PTSD related symptoms at 3 and 6 months, complicated grief at 6 months.

3. **Kindness.** Perception that the team was not kind enough is associated with increased risk of developing anxiety and depression at 3 months, PTSD related symptoms at 3 and 6 months, complicated grief at 6 months.

4. **Preparation for death.** Relatives who were not informed that the patient was dying, who were unable to express important things or to say goodbye were more at risk of developing anxiety and depression at 3 months, PTSD related symptoms at 3, 6 and 12 months and complicated grief at 6 and 12 months.

5. **Presence at time of death is associated with increased risk of developing PTSD related symptoms and complicated grief at 6 and 12 months.**

**Discussion** Relatives are sensitive to interaction between the ICU team and themselves as well as between the team and the patient. Quality of communication (both verbal and non verbal) and support, as well as preparation for the death, are key components of relatives’ experience that impact on grief symptoms in the months that follow the patient’s death.

**Conclusion** Quality of care and support during the dying process are at the heart of the relatives’ experience. This study puts forward practices that may be improved in order to promote both palliative care and family centered care in the ICU and, in fine, decrease grief symptoms in bereaved relatives.

**Competing interests** None.

**O21** Impact of fluid-induced hyperchloremia on acid base balance and outcomes in septic shock: post hoc analysis of the “Hyper2S” study

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Annals of Intensive Care 2017, 7(Suppl 1):O21

Introduction The harmfulness of fluid-induced hyperchloremia (H-Cl) remains debated. Large randomized trial showed that chloride-rich crystalloids did not worsen outcome [1]. The volume of fluids, however, was limited and the incidence of H-Cl was not recorded.

Patients and methods In a post hoc analysis of the database of the RCT “Hyper2S”, a study comparing normal to 3% hypertonic saline for volume of fluids, chloride load (all fluids) was recorded at H0-12-24-72. Episodes of hyperlactatemia (H-Lac > 2 mmol/L), metabolic acidosis (pH < 7.35 + Bic < 22 + Lact > 2) or H-Lac (pH < 7.35 + Bic < 22 + Lact > 2) were recorded. Acute kidney injury (AKI) was defined by doubling creatinine or dialysis.

Results 413 patients without missing data were analysed. H-Cl and HCl-acidosis were recorded in 257 (62%) and 77 (19%) patients, respectively. Baseline severity scores were similar in patients with and without H-Cl but vasopressor dose was higher in patients with H-Cl episode (Table 5).

Mean pH 7.36 (7.29–7.40) 7.32 (7.26–7.37) <0.001
Mean lactate (mmol/L) 137 (79–208) 132 (80–191) 0.31

Evolution from H0 to H72
Volume of fluid resuscitation (L) 1.4 (0.6–2.3) 2.2 (1.1–3.9) <0.001
Chloride load (all fluids) (mmol) 287 (129–491) 690 (293–1127) <0.001
Mean pH 7.36 (7.29–7.40) 7.32 (7.26–7.37) <0.001
≥1 episode metab. acidosis [n (%)] 82 (53) 204 (79) <0.001
≥1 episode H-Cl acidosis [n (%)] NA 77 (30) –
≥1 episode H-Lact acidosis [n (%)] 41 (26) 149 (58) <0.001

Outcomes
AKI [n (%)] 96 (64) 146 (58) 0.42
D28 mortality [n (%)] 60 (38) 95 (37) 0.76

not associated with AKI or mortality (Table 5). The mortality of patients who experienced H-Cl acidosis (25%) was similar to patients who never experienced metabolic acidosis (28%), p = 0.65.

Conclusion H-Cl is frequent in septic shock patients resuscitated with chloride-rich fluids but does not increase AKI or mortality. H-Lact is more frequent in patients with H-Cl, which is an important bias for the interpretation of the origin of acidosis and attributable mortality.

Competing interests None.

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O22
Grief symptoms in relatives of brain dead patients: comparison of relatives of donor and non-donor patients
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Annals of Intensive Care 2017, 7(Suppl 1):O22

Introduction Long after the death of a loved one, end-of-life decisions can remain with the living and have been implicated in post ICU burden. In the case of organ donation, the relatives are at the centre of the decision process; within a limited time frame, they will be told that the patient is brain dead and will be asked to consider organ donation. Attention has been focused on how relatives make the decision to donate or not to donate the patient’s organs but only very few studies have focused on the impact of organ donation decision making on their psychological well-being during the months that follow the patient’s death. The goal of this study was to describe the grieving process of relatives who were approached about organ donation in the context of brain death and to compare grief symptoms of relatives of donor patients (DPs) versus relatives of non-donor patients (NDPs).

Patients and methods We conducted a prospective, observational study in 28 ICUs in France. For each brain dead patient, the relative who served as the surrogate was included at time of organ donation interview: At 1 month, to complete a questionnaire regarding their experience in the ICU; at 3 months to complete the Hospital Anxiety and Depression Scale (HADS) and the Impact of Event Scale-Revised (IES-R) for PTSD symptoms; at 9 months, to complete the IES-R and the Inventory of Complicated Grief (ICG). The primary outcome measure was the IES-R (at 3 and 9 months).

Results 202 relatives were included in the study among which 158 were relatives of DPs and 44 of NDPS. At 1 month after the patient’s death, 79.2% of relatives completed the questionnaire, at 3 months 70.3% completed the HADS and the IES-R and at 9 months 61.4% completed the IES-R and the ICG.
Relatives’ experience of ICU and decision-making varies between the 2 groups. Relatives of NDPs are less satisfied with communication with ICU team than relatives of DPs (26.6 vs 8%, p = 0.021). Relatives of NDPs were more often shocked by organ donation request than relatives of DPs (64.52 vs 19%, p < 0.0001). During organ donation discussions, relatives of NDPs more often declared an absence of support from the ICU team (19.35 vs 1.59%, p = 0.0008) and more often felt under pressure (41.94 vs 7.14%, p < 0.0001).

At 3 months, there was no difference in the IES-R score between relatives of DPs (31 [21–41]) and NDPs (31.5 [11.25–34]) (p = 0.29). Similarly there were no differences in HADS score (13 [9–20] and 13.5 [8.25–20] respectively, p = 1.00) and anxiety and depression subscores. At 9 months, there was no difference in the IES-R score between relatives of DP (26 [12.75–38]) and NDP (33.5 [21.25–43.25]) (p = 0.17). Similarly there were no differences in ICG score (25.5 [14.37–25] and 32.5 [17.43–25] respectively, p = 0.11).

Discussion
Relatives of DPs and NDPs have different experience of quality of communication and quality of support during the patient’s ICU stay. More than the decision itself, quality of the organ donation process impacts on relatives’ grief symptoms.

Conclusion
The decision (donation vs no donation) has no impact on grief symptoms in the months following the patient’s death. However, experience of the request and of the decision itself, as well as quality of communication and support, are elements that effect relatives’ experience and grieving process.

Competing interests
None.

O23
Implementation and impact of the surviving sepsis campaign protocol: results of a quasi-experimental study in Democratic Republic of Congo (DRC)
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Introduction
In developed countries, the application of the “Surviving Sepsis Campaign” (SSC) protocols in the management of sepsis and septic shock has been associated with increased survival [1]. In the context of a fragile health system, the application of these protocols may be expected to prove to be difficult to undertake and the impact may also be expected to vary [2].

Patients and methods
We conducted a prospective, quasi-experimental before and after study in the university hospital of Kinshasa (DRC) comparing 33 consecutive patients in septic shock treated according to usual care versus 39 patients treated according to a protocol type EGDT from 1st February 2014 to 31st July 2014 (pre-protocol phase) and 1st September 2014 to 28th February 2015 (post-protocol phase). Between the 2 phases, we have drawn and implemented a local protocol based on SSC recommendations. A kit consisting of central venous catheters, devices capable of measuring central venous pressure, flow regulator (Dosiflow R) and lactate reader was available.

In the absence of electrically driven syringe pumps, catecholamines were administered by continuous perfusion using a flow regulating device (Dosiflow). We drew up calculating tables based on a standard dilution and a variable administration rate controlled by the flow regulating device, thus permitting precise and constant dose administration of nitroprusside (µg/h).

The main outcome measures were: the rate of compliance with 6-h sepsis care bundles and the hospital mortality until J30. The Student's and U Mann–Whitney tests were used to compare quantitative variables, and the chi-square test to compare qualitative variables.

Kaplan–Meier survival curves were used and the differences between the two curves were analyzed using the log rank test. This study received the approval of ethics committee and was registered 23th January 2014 under number ESP/CE/053/14.

Results
Baseline characteristics of patients were similar in both two groups. Lung infection was the main source of septic shock and antibiotics administration delay was less than 3 h in the two groups. In contrast, blood cultures and blood lactate were performed only in patients of the EGDT group.

Patients of the EGDT group received more IV fluids (+1226 ml on average), more catecholamines (+24, 9%) and were more often transfused (+13, 7%) than the patients treated in the habitual fashion. Protocol compliance was substantially improved, passing from 0 to 50%. The absolute risk reduction of mortality was 17% (100 vs 83%; p = 0.0037) when all the therapeutic measures had been employed.

The Kaplan–Meier survival curves showed that the patients in the pre-protocol group had a significantly greater risk of dying compared with those in the post-protocol group. The average length of stay was 2.2j vs 5.2j p = 0.0037.

Discussion
Our results in terms of compliance with the sepsis bundle were satisfying given that even in highly developed countries the compliances rates are often below 50% [3]. Ignorance of the very existence of these recommendations and logistical factors are often quoted as the reasons why compliance is so poor.

The positive impact that we observed on mortality was found in similar studies [4]. Other more recent randomized controlled studies have not found significant differences but this may be explained by the similarity of treatment protocols in the two groups [5].

Our study is coherent according to current literature but there exist certain limits which might be expected in this kind of study.

The research was implemented in only one centre and these results cannot be extrapolated to all sub-Saharan Africa, nor even to our own country. As in all before/after studies, comparison with a historic control group is liable to introduce a bias based on confusion. We sought to minimize this bias however by choosing a short study period (12 months) with the same therapeutic team during the entire study.

Conclusion
Despite a difficult socio-economic context, the implementation of a local protocol based on the recommendations of the SSC was associated with improved outcome in septic shock patients in our hospital compared to usual care.

We thus intend to continue and to widen the application of this protocol in an effort to evaluate if this positive effect will be observed in a greater number of patients.

Competing interests
None.

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O24 Characteristics and 1-year prognosis of tetanus patients admitted to the ICU
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Introduction
Despite being a fully preventable infectious disease, tetanus is still responsible for about 500,000 death worldwide. In developed countries, the incidence is strongly associated with a lack of vaccination coverage in the elderly. In the USA or in France, people over 65-year-old have a twice to tenfold increase in annual incidence of tetanus versus younger patients [1]. Considering the long lasting effects of tetanus toxin up to 6 weeks, elderly people admitted in intensive care units (ICU) are of particular risk of complications. Prognostics factors are well known in developing countries with a mortality rate above 20% but clinical data in developed countries are missing. We conducted a multi-center retrospective study in France on 90-day and 1-year mortality in patients with tetanus admitted in ICU.

Patients and methods
This study was conducted over 15 French ICUs. All adults patients admitted for tetanus from January 2000 to December 2014 were included. Data were retrospectively collected from medical files. Long-term vital outcome was obtained by interrogating town hall registries.

Results
Seventy patients were recruited over the study period. Median age was 80 years [IQ range 73–84], 86% were women. Median Charlson comorbidity index score was 4 [3–5] and Knaus chronic health score status was distributed equally between category A or B (50% each). All patients had trismus and 56% had generalized form at presentation. The median incubation period was 10 days [IQR 8–14]. Mechanical ventilation (MV) was performed in 90% of patients for a median duration of 36 days [IQR 26–46]. Median SAPS II score at ICU admission was 33 [IQR 26–40], corresponding to an predicted hospital mortality rate of 14%. Median length of stay in ICU was 41 days [IQR 24–53]. Ninety-day and one-year mortality rates were 13% (n = 9) and 16% (n = 11) respectively. Kaplan–Meier survival curve is presented Fig. 4.

Death typically occurred within the first week (55%) due of severe arrhythmia at admission or during the fourth ICU week (withdrawal of life sustaining therapies in 3 patients or multiple organ dysfunction syndrome due to nosocomial infections in 2). Ventilator-associated pneumoniae incidence was 15 episodes per 1000 ventilation-days (total of 2234 ventilator-days observed) corresponding to 48% of ICU patients receiving MV.

Mortality was associated with older age (83 [81–85] versus 79 [73–84] years, p = 0.06) and baclofen use (intrathecal or intravenous, 4/9 non survivors vs. 8/61 in survivors, p = 0.04). Shorter incubation period (under the median delay of 10 days), generalized tetanus, wound debridement (performed in 11 patients) and higher SAPS II (above 30) were significantly associated with longer duration of MV.

Discussion
In-hospital mortality rate was low and consistent with SAPS II estimation (2), despite long-term mechanical ventilation in an elderly population. Surprisingly, baclofen use was associated with an increased risk of death and may be linked to drug-related adverse events. Association of wound debridement with MV duration may either be related to higher bacterial inoculum in larger wound or to per-procedure toxin release.

Conclusion
Long-term follow-up for tetanus-related ICU admission highlights the favorable outcome of this elderly population despite very prolonged MV and frequent infectious complications.

Competing interests
None.

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O25 The clinical spectrum of purpura fulminans in adult patients: a national multicenter retrospective study of 306 patients
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Introduction The available data on Purpura Fulminans in adult patients are scarce, old and mainly limited to patients with meningococcal infections. Our aims were:
1. to describe the clinical features.
2. to identify predictive factors of in-ICU mortality.
3. to compare the presentation and outcomes between causative micro-organisms.
4. to report the rate of limb amputation and identify predictive factors for limb amputation in adult patients admitted in intensive care unit for an infectious Purpura Fulminans.

Patients and methods We performed a 16-year national multicenter retrospective study in 56 ICUs in France from 2000 to 2016. Infectious PF was defined by the association of a sudden and extensive purpura, whatever its causative microorganism, together with the need for vasopressor support. Patients with a noninfectious purpura or with a purpura in a context of infectious endocarditis were excluded from the study.

Results (1) Clinical features upon ICU admission A total of 306 patients were included in the study for an incidence of 0.35 patients per year and per center. Patients were young (median age 34 years [21–53]) and had no previously known comorbidity in 69% of cases. Symptoms before ICU admission included fever (77%), digestive symptoms (61%), headache (45%), myalgia (25%) and lower limb pain (21%), for which 16% of the patients consumed non-steroidal anti-inflammatory drugs before admission. Before ICU admission, 77% of patients had a purpura notified and 75% received a parenteral β-lactam antibiotic. A successful resuscitation of an out-of-hospital cardiac arrest was recorded for 5% of patients. Mean coma Glasgow score at ICU admission was 15 (13–15) and 20% of patients had a neck stiffness. A lumbar puncture was performed in 56% of patients and showed a meningitis (pleocytosis >10/mm³) in 45% of them. A bacteremia was documented in 66% of cases (n = 202/306) and a positive cerebrospinal fluid culture was obtained for 51% of lumbar punctures performed (n = 85/171). In all, the two predominantly identified microorganisms were Neisseria meningitidis (n = 195/306, 63%), mainly serogroup B (39%) and C (34%), and Streptococcus pneumoniae (n = 67/306, 22%).
(2) Patients’ outcomes In-ICU mortality was 41% (n = 126/306). Compared to ICU non-survivors, ICU survivors were younger (29 vs 43 years, p < 0.0001), had more frequently received a β-lactam antibiotic before ICU admission (83 vs. 64%, p < 0.0001), had more frequent neck stiffness (26 vs. 12%, p = 0.004) and cytological meningitis (51 vs. 31%, p = 0.029) and were more frequently documented with Neisseria meningitidis (69 vs. 56%, p = 0.04). Age (OR 1.02/year, 95% CI [1.01–1.04]; p = 0.002) and SOFA score (OR 1.44/point, 95% CI [1.30–1.58]; p < 0.0001) upon ICU admission were the only variables independently associated with ICU mortality.

(3) Impact of the identified microorganism As compared with others, patients with pneumococcal PF were older (p < 0.0001), had more frequent asplenia (48 vs. 2%, p < 0.0001), higher SAPS 2 and SOFA scores together with a higher ICU mortality (52 vs. 36%, p = 0.04), and required more frequent limb amputation (31 vs. 9%, p < 0.0001).

(4) Surviving patients requiring amputations 22% of ICU survivors (n = 39/180) eventually required amputation during their ICU stay with a median of 3 (2–4) limbs amputated. Among ICU survivors (n = 180/306, 59%), those who were amputated (n = 39/180, 22%) were older (42 vs. 26 years, p = 0.016) and presented with higher SOFA and SAPS 2 scores, lower platelets counts (38 vs. 87.10³/mm³, p < 0.0001), more severe kidney injury and higher arterial lactate (7 vs. 5 mmol/L, p < 0.001) and creatine kinase (856 vs. 159 IU/L, p < 0.001) levels than those who were not amputated. By multivariable regression analysis, the following risk factors for limb amputation were identified among ICU survivors: SOFA score upon ICU admission (OR 1.34/point, 95% CI [1.14–1.59]; p < 0.0001), Streptococcus pneumoniae PF (OR 5.05, 95% CI [1.73–14.74]; p = 0.003), and platelets transfusion (OR 6.06, 95% CI [2.20–16.73]; p < 0.0001).

Conclusion Purpura Fulminans is a rare disease mainly affecting young healthy patients. Most patients had an extensive purpura identified before ICU admission and those receiving antibiotics before ICU admission had a lower mortality than others. Neisseria meningitidis and Streptococcus pneumoniae were the main micro-organisms identified. The overall ICU mortality was high and limb amputations were needed in almost one quarter of ICU survivors. Patients with Streptococcus pneumoniae PF had a poorer outcome with a higher ICU mortality and a higher risk of limb amputation.

Competing interests None.

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O26

Cerebral NIRS profiles during premedication for neonatal intubation

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Introduction To date, there is no consensus on which anesthetic protocol should be used before neonatal intubation. As a result, awake intubation, although strongly discouraged by current recommendations, is still common in neonates, especially in France. Many experts consider Propofol, a short-acting anesthetic agent, an appropriate drug for premedication before neonatal intubation. Propofol is known to decrease systemic vascular resistance leading to low arterial blood pressure, which consequently raises concerns about its hemodynamic tolerance. However in neonates, low mean arterial blood pressure (MABP) is poorly correlated with low systemic or cerebral blood flow. NIRS (Near InfraRed Spectroscopy) allows cerebral tissue oxygenation monitoring, reflecting cerebral blood flow. The aim of our study was to compare cerebral oxygenation profiles between a combination of a synthetic opioid plus a muscle-blocker and propofol, used as premedication prior to neonatal endotracheal intubation.

Patients and methods Observational prospective study, conducted in 2 of the 8 centers participating in a randomized, controlled, double-blind, multicenter trial (PRETINTEO study, ClinicalTriial.gov identifier NCT01490580). Patients were randomly assigned (1:1) between “atropine-propofol” and “atropine-atracurium-sufentanil” before elective or semi-urgent intubation in the neonatal intensive care unit. Randomization was stratified on weight and center. Exclusion criteria included low blood pressure defined as a MABP (in mmHg) below gestational age (in weeks). Physiological parameters, including pulse oxygenation (SpO2) and regional cerebral oxygen saturation (rSO2), were collected 1 min prior to induction of anesthesia, and then up to 60 min after. To investigate the balance between oxygen delivery and consumption, cerebral fractional tissue oxygen extraction (FTOE) was calculated as FTOE = (SpO2 – rSO2)/SpO2. The analyzed parameters included changes in rSO2 and FTOE over time and their relative change from baseline value in both treatment groups.

Results From March to August, 2016, 65 neonates were assessed for eligibility. Among them, 28 were finally included in this ancillary study. Their mean (SD) gestational age at birth was 32 (5) weeks. Median [IQR] age and mean (SD) weight at the time of intubation were 0.5 [0.2–5.4] days and 1886 (900) g respectively. At the time this abstract...
was conceived, data validation had not been completed. Results will be available at the time of the meeting.

Discussion To our knowledge, only two studies to date have investigated the cerebral hemodynamic effects of propofol in neonates. Both reported persistent low MABP after a propofol bolus but a very small decrease in rSO2. Only one studied propofol before intubation and suggested that low rSO2 could be attributable to low SpO2 during the procedure, but not to low MABP (Smits et al., J Pediatr, 2016). None of them compared propofol to another anesthetic protocol. Our study is thus the first to compare cerebral oxygenation between two currently acceptable regimens of anesthesia for premedication before neonatal intubation.

Conclusion This observational study is expected to provide useful information about cerebral oxygenation in neonates during intubation and to disentangle the effects of the drugs from the effects of the procedure. We also expect this study will encourage Neonatologists to avoid awake intubation.

Competing interests None.

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O27 A prospective multicentric study of severe cutaneous infections in pediatric intensive care: the SCIPIC cohort
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Introduction Cellulitis (CEL) and Necrotizing Fasciitis (NF) are life-threatening skin infections much more frequent in adults than in children. Therefore, despite major differences in epidemiology, microbiology, and outcome, clinical guidelines for management are directly adapted from adult ICU experience. We designed a prospective multicentric cohort study to describe the clinical course of CEL/NF requiring admission to PICU.

Patients and methods This study was approved by the IRB Paris Nord and the French Data Protection Authority (CCTIRS/ CNIL). After parents’ consent was obtained, thirty centers (France, France Polyneusia, Mayotte and La Réunion Islands, French Caribbs Islands, Belgium, Switzerland, Belgium, Canada, Netherlands, Norway) included on a secure-dedicated-website all patients aged from 28 days to 18 years and admitted to PICU for a suspected or confirmed CEL/NF with association of the three following signs: (i) area of skin inflammation, (ii) rapid evolution or skin necrosis, (iii) variation in core temperature (>38.5°C or <36°C). Premature infants (<37 weeks of gestation) and patients with Stevens Johnson syndrome, Purpura Fulminans or Hereditary Angioedema were excluded. We collected all data concerning past medical history, first clinical signs, PICU stay, antibiotic therapy, microbiology (Staphylococcus and Streptococcus species were send to French Reference Centers), radiological diagnosis, surgery and outcome 1 year after discharge. Data are presented as mean ± SD according to their Gaussian distribution.

Results 50 patients (age: 5.9 ± 5.2 months and weight: 24.2 ± 18.1 kg) were included from October 2011 to April 2016. The diagnosis was suspected 3.5 ± 2.7 days after a known risk factor (46%; surgery: 12; NSAIDS: 10; varicella: 9; cancer: 4). Only two patients had travelled abroad. Skin lesions observed on 10.6 ± 16% of body surface were erythema (28), necrosis (13) and bulla (10) and were located on face (17), legs/feet (16), abdomen/pelvis (14), arms/hands (13) and trunk (10). On admission, mortality and organ dysfunction scores were as follow: PRISM = 9.3 ± 7.5; PIM2 = 12.6 ± 23.3; PELOD D1 = 12.6 ± 3.2 and POPC = 1.6 ± 1.0. Main biological tests showed: hemoglobin: 10.0 ± 2.2 g/dl; neutrophils: 10320 ± 8740/mm3; platelets: 197700 ± 140764 mm3; CRP: 189 ± 122 mg/l; PCT, 69.3 ± 75.3 mg/l; fibrinogen: 5.2 ± 2.2 g/l; ASAT: 108 ± 166 UI/l and ALAT: 61 ± 82 UI/l; proctidemia: 53 ± 13 g/l. 28 patients (56%) showed low blood pressure for age including 14 with oliguria, needing fluid challenges (45 ± 26 ml/kg) and vasopressors for 3 ± 1 days. 29 patients (68%) were ventilated invasively for 0.2 ± 10.5 days, with PEEP = ± 7±3 cmH2O and FiO2 = 58 ± 27% during the first 3 days. 80% of patients received continuous intravenous analgesia (morphine and benzodiazepines).

Half of patients were transfused with blood products and also a half received albumin. While MRI was performed in only 7 cases, CT scan and ultrasound were performed to confirm diagnosis in 23 cases (46%; 14 on day 1) and in 21 cases (42%, 15 on day 1), respectively. 28 patients (56%) were operated. One bacterial strain alone was identified in 36 cases (72%) and at least two in eight cases (16%), including Staphylococcus aureus (17), GASβH-5 (13), Escherichia coli (5), Pseudomonas aeruginosa (4). Antibiotics used were penicillin G (26), clindamycin (21), 3rd generation cephalosporin (12) and rifampicin (9). A third of patients received immunoglobulins and five children (10%) received hyperbaric oxygen (22 sessions). Length of stay in PICU was 12.5 ± 14.7 days. Three patients (6%) died and only two (4%) were readmitted to PICU because of failure directly link to skin lesion. POPC score on discharge was 1.9 ± 1.3. Long term follow-up at 1 year is ongoing.

Discussion This study reports the largest prospective multicentric international cohort of pediatric skin infections needing PICU admission to date. Surgery was the first risk factor reported in our cohort, before varicella. Severe circulatory and respiratory failures on admission in a context of deep biological inflammatory syndrome required aggressive treatments during the first 3 days. Lower part of the body was more frequently involved than reported before in childhood. The diagnosis was quickly suspected and confirmed mainly by ultrasound and CT scan, with the use of MRI still very rare. More than a half of patients have been operated after classical treatments associating immunoglobulins and antitoxin-antibiotics. Infection appeared mainly monobacterial as usually described in children but gram-negative strains are emerging. Mortality was still low and sequella were rare.

Conclusion Cellulitis (CEL) and Necrotizing Fasciitis (NF) are still rare in children with low mortality and organ dysfunction if aggressive treatment including surgery is urgently performed. However, it seems that epidemiology has changed during the last decade in high-income countries. Statistical analysis of this database is ongoing to identify predictors of surgical requirements and to describe more precisely bacterial strains involved.

Competing interests None.

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O28 Immunosuppression induced by septic shock in children: a prospective observational study before a multicenter therapeutic survey. Intensive Care Med. 2015;41(8):1506–8.

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Introduction Immunosuppression induced by sepsis is well described in adults. Therapeutic trials with immunomodulatory treatments are already underway, using HLA-DR expression on monocytes (mHLA-DR) or lymphopenia as biomarkers of immunosuppression. Pediatric patients with Septic Shock (SS) have been much less studied. Thus, the main objective of this study was to explore post-sepsis immunoparalysis in a pediatric cohort study. Both sides of cellular immunity were assessed: innate immunity (mHLA-DR) and adaptive immunity (lymphocyte subsets). We also wanted to obtain normal values of mHLA-DR in healthy children, according to age.

Materials and methods

We performed a single-center prospective study with children under 18 years-old, admitted in Pediatric Intensive Care Unit for SS (“Surviving Sepsis Campaign” criteria), between September 2014 and July 2016. We recruited controls from healthy children hospitalized for an elective surgery, without any criteria of infection. mHLA-DR, total lymphocyte count, and lymphocyte sub-populations’ proportions (CD4+ and CD8+ T cells, regulatory T cells, NK cells, and B cells) were determined by flow cytometry. Samples were analyzed at Day 1, 3 and 7, after sepsis onset. Clinical data were collected prospectively, especially severity scores [PIM2, PELOD2 and Cumulative Vasopressor Index (CVI)], and secondary nosocomial infection occurrence.

Results

30 controls and 26 patients were recruited. mHLA-DR in healthy children presented no variation according to age, and was similar to healthy adults. At each time points, mHLA-DR in SS group was decreased, comparing with controls. The medians of mHLA-DR were respectively 6.066 IQR [3.737–16.310], 6.308 IQR [3.185–8.965] and 9.323 IQR [6.384–12.738], versus 29.668 ab/c IQR [24.335–39.199] in the control group (p < 0.001 at D1, D3 and D7; Mann–Whitney). mHLA-DR at D3 was significantly correlated with the Cumulative Vasopressor Index (Spearman’s correlation coefficient r = −0.50; p = 0.031). Patients secondarily infected presented a lower mHLA-DR than patients without secondary infection; respectively 4398 ab/c [2437–6212] versus 8474 ab/c [5904–10844] (p = 0.022, Student t test) (Fig. 5).

Total lymphocytes and CD4 T cells were decreased at D1 and D3. This lymphopenia corrected between D3 and D7. NK cells were decreased at each time points. B lymphocytes were decreased comparing to controls at D1 and D7 (not D3), but much less pronounced than T cells. Regulatory T cells percentage was initially comparable to controls. Then, a gradual increase was observed, which became statistically significant at D7, compared to controls (p = 0.0058; Mann–Whitney).

No significant difference was observed between patients with or without secondary infections, according to any lymphocyte subsets.
screen VAP. However, sensitivity should be improved by adapting VAE criteria to children.

**Conclusion** Using the CDC updated VAP definition for children, VAP incidence is similar to adults. Adult VAE cannot be used to screen prospectively VAP in children. Specific pediatric VAE needs to be developed and validated.

**Competing interests**

None.

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**O30 Incidence and risk factors of ventilator associated pneumonia in neonatal intensive care unit**

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**Annals of Intensive Care 2017, 7(Suppl 1):O30**

**Introduction** Ventilator associated pneumonia (VAP) is defined as a lung infection occurring after 48 h of mechanical ventilation. Ventilator associated pneumonia incidence, complications and mortality are well known in critically ill adults but probably under diagnosed in critically ill neonates. The main objective of our study was to evaluate the incidence of VAP in neonatal intensive care units, evaluate associated mortality and morbidity and find potential risk factors.

**Patients and methods** This is a prospective, observational, single-center conducted in neonatal intensive care unit of Armand Trousseau hospital from 01/11/2014 to 31/10/15. All infants aged 28 days or less hospitalized in the service are included.

**Results** 381 patients were enrolled including 327 intubated patients. 17 of 327 patients intubated presented VAP. The incidence of VAP was 4.05 per 1000 days of invasive ventilation with an incidence rate of 8.78 per 1 000 days of invasive ventilation. The average age at diagnosis of VAP was 21.47 ± 13.02 days for an average duration of invasive ventilation of 15.23 ± 11.48 days. After VAP, invasive ventilation time is prolonged to 25.71 ± 20.90 days (OR 1.19 [1.12–1.27]). The nosocomial infection rate was significantly higher in the VAP group (p < 0.001) with 9 on 17 patients in VAP group (52.94%) versus 21 on 310 patients (6.77%). The occurrence of VAP was significantly associated with higher mortality (OR 4.46 [1.32–14.94]) and an increase in invasive ventilation times (p < 0.001) and non-invasive (p < 0.001) and hospital stay (p < 0.001). There is a significant difference in the duration of invasive ventilation before VAP [average 15.23 ± 11.48 (1–35)] compared to patients without VAP [mean 4.84 ± 4.92 (1–28), p < 0.001]. Patients with a birth weight less than or equal to 1000 grams are associated with risk of VAP (OR 4.31 [1.38–13.39]) in multivariate analysis contrary to the term. Intubated patients with a balloon are associated with risk of VAP (adjusted OR 4.03 [1.14–14.26]) and a Snappe-II score above 16 also (OR 4.98 [1.40–17.67]).

**Conclusion** We managed one of the larger neonatal study for VAP in critically ill neonates. These pneumonias remained frequent in critically ill neonate and are associated with a higher mortality and morbidity. Patients with a birth weight less than 1000 g seem to be particularly vulnerable.

**Competing interests**

None.

**O31 Extracorporeal life support for acute respiratory failure in immunocompromised patients: an international multicenter retrospective study (The IDEA study)**

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**Introduction** The proportion of immunocompromised patients with extracorporeal life support (ECLS)-treated severe ARDS varies from 5 to 31% in recent cohorts. To date, very few data on ECMO use and its associated outcomes are available on this population. Our aims were to (1) describe the clinical features, (2) compare the outcomes between causative immunocompromised status, (3) identify predictive pre-ECLS factors of 6-month mortality, and (4) report the rate of ECLS-related complications.

**Patients and methods** We performed an international multicenter retrospective study in 10 ICUs from 2008 to 2014. Immunocompromised status was defined by hematologic malignancies, solid tumor, solid organ transplant, human immunodeficiency virus (HIV), long-term or high dose glucocorticoids or immunosuppressant use. Inclusion criteria were immunocompromised patient with acute respiratory failure rescued by extracorporeal membrane oxygenation or (ECMO) or extracorporeal CO2 removal (ECCO2R).

**Results** 1. A total of 225 patients (age 48.6 ± 15.1 years; APACHE II score 28% were included in the study (30% hematologic malignancies, 28% long-term corticosteroids or immunosuppressant use, 18% solid tumor, 16% solid organ transplant, and 9% HIV). ECLS was initiated for severe ARDS, moderate ARDS, or chronic end-stage respiratory in 190(84%), 14(6%), and 19(8%) patients, respectively. Main ARDS etiologies were bacterial pneumonia (30%), viral pneumonia (18%), and specific lung involvement (12%). Refractory hypoxemia (73%) was the main indication for ECLS with lowest pre-ECLS PaO2/FiO2 at 63 (51–87) mmHg. Venovenous ECMO and ECCO2R were initiated in 199 (88%) and 15(7%) patients, respectively. Pre-ECLS tidal volume was 5.7(4.7–6.4) ml/kg, with a positive end-expiratory pressure at 10(8–13) cmH2O and a plateau pressure at 32(30–35) cmH2O. Interval between mechanical ventilation onset and cannulation was 2(1–8) days.

2. Six-month mortality of patients with hematologic malignancies, long-term corticosteroids or immunosuppressant use, solid tumor, solid organ transplant, and HIV were 76, 60, 80, 51, and 71%, respectively (log-rank test, p = 0.002). Cumulative survival at 6 months were lower for patients with hematologic malignancies versus others (log-rank test, p = 0.003) whereas those with solid organ transplant exhibited higher cumulative survival at 6 months (log-rank test, p = 0.02).

3. One hundred and three patients (46%) were successfully weaned from ECLS. In-ICU and 6-month post ICU discharge survival were 37% (n = 83/225) and 32% (n = 72/225), respectively. Compared to patients who died within 6 months after ICU admission, 6-month survivors were younger (46 vs. 49 years, p = 0.05), had more frequently a newly diagnosed immunocompromised status (39 vs. 17%, p = 0.003), were more frequently patients with solid organ transplant (24 vs 12%, p = 0.02, had a higher pre-ECMO hemoglobin (9.4 vs 8.8 g/dL) and platelet counts (160 vs 112 × 10^3/µL; p = 0.008), and exhibited lower mechanical ventilation-ECMO onset interval (10–5) vs 3(1–9) days, p = 0.002). Age (OR 1.02/year, 95% CI [1.002–1.04], p = 0.035), solid organ transplant (0.38 [0.17–0.85], p = 0.019), newly diagnosed immunocompromised status (0.32 [0.16–0.65], p = 0.002), platelet count ≥200,000 × 10^3/µL (0.33[0.15–0.72], p = 0.005) and delay from mechanical ventilation initiation to ECMO cannulation >7 days (3.23
Driving pressure is a significant predictor of mortality in the acurasy and proseva randomized controlled trials in ARDS patients

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Introduction Driving pressure (ΔP) across the respiratory system has been suggested as the strongest predictor of hospital mortality in ARDS patients. We wonder whether this result may be due to the wide range of tidal volume (VT) and PEEP used across the trials included and whether a strict control of them would minimize the role of ΔP as predictor. Therefore, we investigated ΔP in two trials in which lung protective mechanical ventilation was applied to ARDS patients. Our working hypothesis was that ΔP was a risk factor just like compliance (Crs) or Plateau pressure (Pplat) of the respiratory system.

Patients and methods ARDS patients included in the Acurasy and Proseva trials previously reported were used. Both had inclusion criteria (notably PaO2/FIO2 < 150 mmHg and PEEP ≥ 5 cm H2O) and similar lung protective mechanical ventilation (in particular VT 6 ml/kg predicted body weight and PEEP/FIO2 table). Both found survival benefit in the experimental group. SOFA, continuous neuromuscular blocking agent (NMBA) infusion, prone position, combined use of NMBA and prone position, pH, PaCO2, PaO2/FIO2, lactate, breathing frequency, VT, PEEP, Pplat, Crs and ΔP were recorded at day 1 after inclusion together with gender, age and SAPSII at the time of admission and compared between survivors and nonsurvivors at day 90. Cox proportional hazard models were used with covariates significantly different between survivors and non survivors at the threshold of 0.20 and mortality at day 90 as dependent variable. Due to the obvious colinearity between ΔP, Crs and Pplat we performed the following analyses. First we made a specific Cox model for each of them. Second, we developed three Cox models in which we used the above variables by couples (Pplat-ΔP; Crs-ΔP and Crs-Pplat). We made the following assumptions: if both variables in the couple lacked significance in the second model, the same information was carried by each component of the couple; if one or both variables kept significant correlation each brought significant and distinct information; if significant correlation was kept for one of the variables in the couple and lost for the other the former would be more informative than the latter.

Results Both trials enrolled 805 patients of whom 787 had data available at day 1 of whom 533 survived and 254 did not. In the univariate analysis, ΔP averaged 13.7 ± 3.7 and 12.8 ± 3.7 cmH2O (P = 0.002) in nonsurvivors and survivors, respectively. In each single Cox model, Hazard ratios (HR) were 1.05 (1.02–1.08) (P = 0.005), 1.04 (1.01–1.08) and 0.985 (0.972–0.999) (P = 0.023) for ΔP, Pplat and Crs, respectively. PEEP and VT were not significant risk factors in any model. In the model with ΔP and Pplat used together, each of them kept significance [HR 1.31 (1.07–1.61) and 1.13 (1.02–1.26)]. In the model with ΔP and Crs both lost significance and the same was true in the model using Pplat and Crs.

Discussion ΔP was a significant predictor with a 5% increase in mortality per each cmH2O increment of ΔP, a result similar to that found by Amato et al. However, Pplat conveys the same information. This is likely due to the fact that VT and PEEP were similar in each group of both trials and not different between survivors and non survivors.

Conclusion When strict lung protective mechanical ventilation is applied to ARDS patients ΔP, Crs and Pplat were risk factors of mortality. ΔP and Pplat bring distinct and significant information.

Competing interests None.

O33 Impact of PEEP and body inclination in the supine and prone positions on esophageal pressure in ARDS patients

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Introduction Prone positioning (PP) for long sessions in association with use of low tidal volume and cisatracurium infusion have been shown survival benefit in ARDS patients. Little is known about evaluation of PEEP setting based on end-expiratory transpulmonary pressure at zero flow (PEEPtotL) in PP. We hypothesized that esophageal pressure (Pes) was lower in PP than in supine position due the relief of the weight of the mediastinum. Hence, the PEEP level which lung protective mechanical ventilation was applied to ARDS patients ΔP, Crs and Pplat were risk factors of mortality. ΔP and Pplat bring distinct and significant information.

Competing interests None.

Materials and methods A prospective interventional physiologic study was performed in patients with severe ARDS (PaO2/FIO2 < 150) and requiring PP. Pes was measured with an esophageal balloon. Transpulmonary pressure (Ptp) was computed as the difference between airway pressure and Pes. DPL was computed as the difference in Ptp at the end of inspiration and at the end of expiration at zero flow. Chest wall elastance (E(st,cw)) was calculated as the ratio of end-inspiratory Pes minus PEEPtotL divided by tidal volume. End-expiratory lung volume (EELV) was measured by nitrogen wash-out method. Thorax angulation was 30° and 0° in supine position (SP) and 0° and 15° in PP. From PEEP totL adjusted according to the low PEEP/FIO2 table of the ARMA trial (1), PEEP level was further adjusted to achieve PEEPtotL close to 3 cmH2O. Measurements were done in supine position and after 1 h in PP.

Results Twenty patients were included. PEEPtotL did not significantly vary between SP(30°) and PP(0° or 15°). However, opposite variations were found according to thorax angulation in PP: a rise of 1.9 (SD 1.7) cmH2O in PP(0°), p = 0.005 and a drop in PP(15°) of 2.2 (SD 3.2) cmH2O, p = 0.06. In a complementary analysis, PEEPtotL was studied in four positions (SP(30°), SP(0°), PP(0°) and PP(15°)) and followed an inverted U-shape pattern with mean values of 8.6 (SD 3.2), 12.8 (SD 2.7), 10.2 (SD 3.3) and 2.6 (SD 2.1) cmH2O, respectively. These differences were statistically significant (Holm adjusted p value for multiple comparisons <0.05) except between SP(30°) and PP(15°).
As a consequence, PEEPtotL rose of 2.6 cmH2O between SP(0°) and PP(0°). With postural variations, EELV was significantly altered. Hence, PEEPtotL at PP(0°) was computed at the EELV in SP(0°), allowing to eliminate the impact of the change of EELV between postures: 77% of PEEPtotL changes were due to change of posture per se [10.4 (SD 3) vs. 8.5 (SD 3.2) L/cmH2O, \( p = 0.004 \)]. Chest wall elastance did not change between 30° supine and 0°PP [10.4 (SD 3) vs. 8.5 (SD 3.2) L/cmH2O, \( p = 0.85 \)].

Discussion Variation of thorax angulation significantly alters Pes and Ptp values. Splitting the variation of Pes into those due to change in either EELV or posture allows a better understanding of the impact of PP on Pes.

Conclusion PP(0°) was associated with Pes rise as compared with SP(30°) and Pes drop when compared with SP(0°). However, when analyzing PEEPtotL variation due only to posture change (and not EELV change), a rise of almost 3 cmH2O of PEEPtotL was seen in PP(0°) compared to SP(0°) or SP(30°). Clinicians might therefore consider lowering PEEP level in PP.

Competing interests None.

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O34 Can we consider criteria for acute respiratory distress syndrome (ARDS) in patients breathing spontaneously?
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Introduction According to the Berlin definition for ARDS, the degree of hypoxemia must be assessed under either invasive or noninvasive ventilation (NIV). In a recent large international survey, about 15% of ARDS were diagnosed while treated with NIV. However, NIV is debated in hypoxemic patients with acute respiratory failure. We aimed to assess whether the use of NIV is really needed to diagnose ARDS in patients with spontaneous breathing under oxygen.

Patients and Methods We included all ICU patients treated first with standard oxygen and then with NIV for non-hypercapnic acute respiratory failure from 2 prospective studies [1, 2]. \( \text{PaO}_2/\text{FiO}_2 \) was assessed at ICU admission under oxygen with an easily bedside calculated \( \text{FiO}_2 \), and under NIV after 1 h of initiation and within the first 24 h. Severity of hypoxemia was considered as mild when \( \text{PaO}_2/\text{FiO}_2 \) ranged from 201 to 300, moderate from 101 to 200 and severe when \( <100 \text{mmHg} \).

Results Among the 219 patients with acute respiratory failure treated with NIV, 172 (79%) fulfilled ARDS criteria within the first 24 h following ICU admission. ARDS was classified as mild in 14%, moderate in 53%, and severe in 33% of cases. The overall rates of intubation and ICU mortality were 58 and 30% respectively.

When considering the 155 patients with bilateral infiltrates on chest X-ray and \( \text{PaO}_2/\text{FiO}_2 < 300 \text{mmHg} \) under oxygen, 87% (\( n = 135 \)) had ARDS criteria after 1 h of NIV and 95% (\( n = 148 \)) within the first 24 h.

Conclusion Almost all patients with bilateral infiltrates and \( \text{PaO}_2/\text{FiO}_2 \leq 300 \text{mmHg} \) under oxygen standard meet ARDS criteria after NIV initiation. Mechanical ventilation does not seem necessary to diagnose ARDS in patients with spontaneous breathing.

Competing interests None.

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O35 Prognosis factors of severe influenza in ICU and introduction delay of oseltamivir
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Introduction Influenza infection has a major impact on ICU hospitalizations during epidemic season, mainly affecting the elderly with exacerbation of their comorbidities. To identify bad prognosis factors in severe influenza and the impact of delayed antiviral therapy among ICU patients infected with influenza.

Patients and methods We conducted a retrospective, observational study in Angers CHU medical ICU, from November 2009 to April 2015. Influenza infection was confirmed by Immunofluorescence. Demographic data, comorbid conditions, antiviral treatment and time elapsed between ICU admission and treatment, clinical outcome, bacterial infection associated, type of virus, antibiotic, ventilation, renal injury, use of vasopressive drugs were recorded. Risk factors for death were analyzed by backward stepwise logistic regression.

Results The study population consisted of 85 patients whose mean age was 63 years old. 80% had at least one comorbidity. Influenza A infected 71% of our patients and type B 28%. [iz1] Most of them was admitted for respiratory distress and 59% were under invasive ventilation. The rate of death was 25% at day 28. 40% were under vasopressive drug. 84% received NAI among whom 76% received it in the first 2 days after ICU admission. Risk factors for death were shock at the admission, immunosuppressive conditions and late administration of NAI. Patients receiving NAI more than 48 h after ICU admission had a higher risk of death.

A multivariate analysis was performed. Shock at the admission was associated with mortality (OR = 2.38: 95% CI 0.03–0.045) and time elapsed from ICU to [RM1] NAI less than 3 days was associated with mortality (OR −2.38: 95% CI 0.03–0.045) and time elapsed from ICU to [RM1] NAI.

See Table 6.

Conclusion Cases of severely ill suspected influenza infections in epidemic period may benefit as soon as they are admitted in ICU from antiviral therapy. Our results add to the existing observational data on the effectiveness of starting antiviral treatment with Neuraminidase Inhibitors as soon as Influenza is clinically suspected, even if suspicion appears at ICU admission and with delay from symptoms onset.

Competing interests None.
## Table 6  See text for description

|                  | Death d28 | Alive d28 | p      |
|------------------|-----------|-----------|--------|
| **Baseline characteristics** |           |           |        |
| Age              | 66.1 ± 12.8 | 61.9 ± 17.2 | 0.3    |
| Sex (M)          | 12 (54.5) | 28 (45.2) | 0.45   |
| BMI              | 25.5 ± 5.3 | 28.1 ± 8.6 | 0.2    |
| Corticosteroids in 10th days before ICU admission | 5 (23.8) | 5 (7.9) | 0.052  |
| **Clinical presentation** |           |           |        |
| Charlson score   | 3.2 ± 3.4 | 2.1 ± 2.5 | 0.16   |
| IGS II           | 49.1 ± 20.8 | 36.5 ± 15.5 | 0.0039 |
| **Clinical complications** |           |           |        |
| Fever (>38 °C)   | 18 (85.7) | 40 (63.5) | 0.056  |
| Body aches       | 1 (4.8)   | 16 (25.4) | 0.085  |
| **Initial influenza diagnostic test** | 0.77 |          |        |
| Rapid diagnosis  | 8 (38.1)  | 26 (41.3) |        |
| Immunofluorescence | 11 (52.4) | 28 (44.4) |        |
| **Clinical complications** |           |           |        |
| Shock            | 18 (85.7) | 16 (25.4) | <0.0001 |
| ARDS             | 11 (52.4) | 0 (0.0)   | 0.047  |
| Acute renal failure | 12 (57.1) | 13 (20.6) | 0.0015 |
| Secondary bacterial infection | 13 (61.9) | 22 (34.9) | 0.03   |
| **Details of antiviral treatment** |           |           |        |
| ICU to Neuramin delay <3 days | 9 (42.9) | 45 (71.4) | 0.018  |

At the same time, potassium level decreased slightly in the placebo arm (p < 0.01), when glucose and lactate levels increased (NS). Symmetric changes were observed in the salbutamol arm after the second nebulization (+13.7 vs. −8.9%, p < 0.01). Nebulization was well tolerated in all patients.

**Conclusion** Placing the nebulizer immediately after the Y-piece is appropriate to deliver aerosols in NIV circuit and compatible with crimped circuit. Salbutamol nebulization is well tolerated during NIV and increase significantly FEV1 when compared to placebo. If this physiological effect is associated with clinical benefit remains to demonstrate.

### Competing interests
I, the undersigned, certify that the BANNISTER clinical trial «Beta Agonist Nebulization in Non Invasively ventilated Chronic Obstructive Pulmonary Disease (COPD) patients: Safety and Therapeutic Efficacy Range», was funded by Association pour la Promotion à Tours de la Réanimation Médicale and the bench study by Inserm. The nebulizers were provided by Aerogen; the company was not involved in study design, data exploitation or drafting of the manuscript.

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### O36
**Salbutamol nebulization during non-invasive ventilation in exacerbated COPD patient**

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**Introduction** Non invasive ventilation (NIV) is a widely used technique to treat hypercapnic acute respiratory failure (ARF) in Chronic Obstructive Pulmonary Disease (COPD) patients. Nebulization of beta 2 agonists is recommended to treat COPD exacerbations. Nebulizing beta 2 agonist on NIV circuits could be potentially helpful but had been studied only in stable COPD patients. Our aim was to compare effectiveness of salbutamol and placebo nebulized in NIV circuit in patients admitted to the intensive care unit (ICU) for exacerbation of COPD. Aerosols were generated by a vibrating mesh nebulizer, whose optimal place in the circuit (just after the Y-piece) was determined by a preliminary bench study.

**Patients and methods** We conducted a double-blinded trial comparing salbutamol and placebo delivered in a random order as two aerosols separated by 60 min, during NIV, in 43 patients admitted to the ICU for ARF. A spirometry was obtained at several points before and after nebulizations, as were clinical and biological safety parameters.

**Results** Forced expiratory volume in one second (FEV1) increased significantly from baseline to 40 min after the end of salbutamol nebulization, when compared to placebo (+5.6 vs. −8.9%, p = 0.04).

**At the same time, potassium level decreased slightly in the placebo arm (p < 0.01), when glucose and lactate levels increased (NS). Symmetric changes were observed in the salbutamol arm after the second nebulization (+13.7 vs. −8.9%, p < 0.01). Nebulization was well tolerated in all patients.**

**Conclusion** Placing the nebulizer immediately after the Y-piece is appropriate to deliver aerosols in NIV circuit and compatible with crimped circuit. Salbutamol nebulization is well tolerated during NIV and increase significantly FEV1 when compared to placebo. If this physiological effect is associated with clinical benefit remains to demonstrate.

### Competing interests
I, the undersigned, certify that the BANNISTER clinical trial «Beta Agonist Nebulization in Non Invasively ventilated Chronic Obstructive Pulmonary Disease (COPD) patients: Safety and Therapeutic Efficacy Range», was funded by Association pour la Promotion à Tours de la Réanimation Médicale and the bench study by Inserm. The nebulizers were provided by Aerogen; the company was not involved in study design, data exploitation or drafting of the manuscript.

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### O37
**Effect of a musical intervention on tolerance and efficacy of non-invasive ventilation: the ‘MUSique pour l’Insuffisance Respiratoire Aiguë—MUS-IRA’ randomized controlled trial**

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**Annals of Intensive Care 2017, 7(Suppl 1):O37**

**Introduction** One of major determinants of non-invasive ventilation (NIV) success is its tolerance. Numerous strategies have been assessed to improve this tolerance, such as pharmacological sedation or sophrology. Music therapy has been showed to be effective in healthcare settings, in particular in ICU invasively ventilated subjects [1]. We therefore investigated the effect of a musical session provided by a trained caregiver on NIV tolerance and efficacy in ICU patients with acute respiratory failure.

**Patients and methods** MUS-IRA is a randomized 3-centers, 3-arm open-label trial (PHRIP 2013). Subjects included were adult acute respiratory failure patients for whom the physician in charge considers NIV as indicated with a level of consciousness allowing a benefit from NIV as indicated with a level of consciousness allowing a benefit from NIV with acute respiratory failure. NIV was conducted in the same fashion in the three arms of randomization. The “musical intervention” subjects received a 30-min L-type music session with the MUSIC CARE© software (2), with
a sleeping mask concealing the eyes. The “sensory deprivation” group had the sleeping mask and the insulating around-ear headphone, during a 30-min period. The “NIV alone” group had their NIV conducted as it is usually in our ICUs. The main objective was to determine if a musical intervention improved NIV tolerance, measured by respiratory comfort, and ventilation parameters at 30 min of NIV in comparison to conventional care. The respiratory comfort was assessed by a nurse or nurse assistant blinded to the treatment arm with a numeric visual scale (from 0 to 10) at the initiation (T0), after 30 min (T30), and at different time points of each NIV session. The primary endpoint was the change in respiratory comfort at initiation and after 30 min of the first NIV session. The secondary endpoints were NIV failure and the percentage of patients requiring anxiolytics or sedative during NIV sessions. The comparison performed in a pre-specified fashion was between “musical intervention” group and “NIV only” group. A total of 99 subjects had to be randomized in order to show a 2-unit difference in respiratory comfort between two groups. This number was extended to 114 because of missing data on the primary endpoint.

**Results** Among the 114 subjects randomized (May 2015 to May 2016)—“musical intervention” group n = 37; “sensory deprivation” group n = 38, “NIV only” group n = 39), median age was 67 years (60–74), and 63 were men (55.3%). Mean baseline respiratory comfort for “musical intervention,” “sensory deprivation” and “NIV only” was 4.34 ± 3.01, 4.24 ± 2.59, 3.89 ± 3.03 respectively for the first NIV session (p = 0.74). Mean change in respiratory comfort between T0 and T30 for “musical intervention”, “sensory deprivation” and “NIV only” was respectively 0.54 ± 3.57; 0.55 ± 2.33; 0.66 ± 2.9. The comparison between “musical intervention” and NIV-alone group yielded a p-value of 0.91. NIV failure during ICU stay was evidenced in 16.2%, 10.5%, 12.8% (“musical intervention”, “sensory deprivation” and “NIV only”, respectively, p = 0.74). No subjects required the administration of anxiolytics or sedatives to cope with the first NIV session in all groups.

**Discussion** Our study failed to evidence a significant effect of a musical intervention on the reduction of respiratory discomfort during the first NIV session of acute respiratory failure ICU patients. One may hypothesize that the improvement provided by NIV (which patients received in all three groups) was such that it outweighed and/or masked a potential effect of the musical intervention. Further analysis are planned to investigate the respiratory comfort and the evolution of physiological parameters during the subsequent sessions.

**Conclusion** Applying a musical intervention early in the course of a treatment by NIV during acute respiratory failure did not modify respiratory comfort. Further research is warranted.

**Competing interests** None.

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**O38**

Apnoeic oxygenation via high-flow nasal oxygen combined with non-invasive ventilation preoxygenation for intubation in hypoxaemic patients in intensive care unit: the randomised OPTINIV study

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**Annals of Intensive Care 2017, 7(Suppl 1):O38**

**Introduction**

Tracheal intubation in the intensive care unit (ICU) is associated with severe life-threatening complications including severe hypoxemia [1]. Preoxygenation before intubation has been recommended in order to decrease such complications. Noninvasive ventilation (NIV)-assisted preoxygenation allows increased oxygen saturation during intubation procedure [2]. However, the NIV-mask has to be taken off after preoxygenation to allow the passage of the tube through the mouth. High-flow nasal cannula oxygen (HFNC) has a potential of apneic oxygenation during the apneic period following the preoxygenation with NIV. We hypothesized that application of HFNC combined with NIV was more effective at reducing oxygen desaturation during the intubation procedure compared to NIV alone for preoxygenation in hypoxemic ICU patients with acute respiratory failure.

**Patients and methods** We did this randomised, controlled, single-center trial with assessor-blinded outcome assessment in patients admitted to ICU. Hypoxaemic patients requiring orotracheal intuba- tion for respiratory failure were randomised to receive preoxy- genation using HFNC [flow = 60 L/min, fraction of inspired oxygen (FIO2) = 100%] combined with NIV (pressure support = 10 cmH2O, positive end-expiratory pressure = 5 cmH2O, FIO2 = 100%) in the interventional-group or using NIV alone in the reference-group (Fig. 6). The primary outcome was the minimal oxygen saturation (SpO2) during the intubation procedure. Secondary outcomes were intuba- tion-related complications and ICU-mortality. The OPTINIV trial was registered with ClinicalTrials.gov, number NCT02530957.

**Results** Between July-2015 and February-2016, we randomly assigned 25 and 24 patients in the interventional and reference groups. During the intubation procedure, median minimal SpO2 values were significantly higher in the interventional-group when compared with the reference-group (100 [95–100] vs 96 [92–99]%, p = 0.029). After exclusion from analysis of two patients for protocol violation, no (0%) patients in the interventional-group and five (21%) patients in the reference-group had a SpO2 below 80% (p = 0.050). We recorded no significant difference in intubation-related complications and ICU-mortality between groups.

**Conclusion** A novel strategy of preoxygenation adding apneic high-flow oxygen to NIV to perform orotracheal intubation in hypoxaemic patients is more effective at reducing oxygen desaturation in comparison to the reference-method using NIV alone.

**Competing interests** None.

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![Fig. 6](image_url)
Feasibility and validity of an observational scale as a surrogate of dyspnea in non-communicating intubated patients in the intensive care unit (ICU)

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Introduction Dyspnea is common and severe in intensive care unit (ICU) patients. In these patients, dyspnea is associated with a poorer outcome. The measurement of dyspnea involves a self-assessment by the patient with a visual analog scale (VAS-Dyspnea), which by definition requires a certain level of communication. However, many intubated critically-ill patients are unable to reach this level of communication, which makes very difficult the assessment of dyspnea.

Recently, our team has developed, a behavioral score named IC-RDOS [1], which is reliable to predict severe dyspnea in ICU patients. This score has been only validated in non-intubated conscious ICU patients. The objective of our study was to validate this score in intubated and non-communicating (NC) patients. IC-RDOS was compared to VAS-Dyspnea in communicating patients and to electrophysiologic markers of dyspnea such as surface electromyographic (EMG) activity of extra diaphragmatic inspiratory muscles [2], in NC patients.

Materials and methods Between February and August 2016, invasively mechanically ventilated patients with clinical respiratory discomfort were included. The IC-RDOS and EMG activity of the Parasternal and Alae Nasi were measured before (baseline, BL) and after one or two interventions aiming at reducing dyspnea. These interventions consisted in an optimization of ventilator settings (Opt) followed, if necessary, by the injection of opioids. The VAS-Dyspnea was only recorded in communicating patients. Non-communicating state was defined by the impossibility to perform a VAS-Dyspnea. This work was supported by the "Fondation de dotation Recherche en Santé Respiratoire" and the "Fondation du Souffle".

Results 35 patients (age 67 [56–76], SAPS II 51 [35–61]; med [IQR] were included; 17 were NC. Among the 18 communicating patients we observed a strong correlation between the IC-RDOS and the VAS-Dyspnea (rho = 0.545, p = 0.038). Among NC patients, IC-RDOS decreased from 6.2 [4.3–6.6] at BL to 4.4 [2.7–6.3] (p = 0.286) and to 4.3 [2.9–4.8] (p = 0.024) after Opt and opioids injection, respectively. Among NC patients, we observed concomitantly, a significant diminution of the EMG activity (EMGAUC) of the Alae Nasi compared to BL (−38% [p = 0.003] and −64% [p = 0.008] after Opt and morphine injection respectively) and a significant diminution of the EMGAUC of the Parasternal relative to BL (−65% [p = 0.039]) after morphine injection. Among NC patients, we found a significant positive correlation between IC-RDOS and Parasternal EMGAUC (rho = 0.538, p = 0.060).

Conclusion IC-RDOS is a reliable surrogate of dyspnea in communicating and non-communicating intubated ICU patients.

Competing interests None.

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Prevalence and impact of pleural effusion during weaning from mechanical ventilation: a prospective multicenter study

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Introduction Pleural effusions are common in critically ill patients. Bedside lung ultrasonography (LU) outperforms chest radiograph for the diagnosis of pleural effusion. Pleural effusion drainage improves oxygenation and respiratory mechanics of patients under mechanical ventilation. However, the role of pleural effusion during the weaning process is unclear.

Patients and methods We performed a prospective multicenter study in six ICUs in France. We used ultrasonography to screen for pleural effusion during the weaning process. Weaning failure was defined as failure of spontaneous breathing trial (SBT) or extubation. Patients were included the day of a first weaning test after at least 24 h of mechanical ventilation. We evaluated associated factors with pleural effusion and its evolution.

Results Two hundred and forty nine patients were included. Median duration of mechanical ventilation was 4 [2–8] days before a first SBT. Forty-seven patients (19%) failed a first SBT and 31 (15%) had extubation failure (reintubation or death within the 7 days following extubation). Pleural effusion was detected in 139 patients (56%) the day of SBT. Most of pleural effusions were homogeneously anechoic (98%) and associated with pulmonary condensation or atelectasis (70%). Interpleural distance was higher among patients with left ventricular diastolic dysfunction, cancer history, ARDS or dialysis before the first SBT. A higher interpleural distance was associated with SBT failure and weaning failure. Patient with weaning failure had more often large pleural effusion (34 vs 12%, p < 0.01). Among patient failing the first SBT and followed up during the weaning process, either diuretic nor fluid balance change altered the interpleural distance within 24–48 h.

Conclusion Pleural effusion is frequent during the weaning process and is associated with weaning failure. Further studies are needed to test whether a strategy aimed at draining large pleural effusions has the potential to decrease duration of weaning process.

Competing interests None.

Characterization of emergency physicians’ recourse to intensive care unit physicians: a prospective multicenter study

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Introduction Recourses to the intensive care units (ICU) physicians by the emergency department (ED) physicians are frequent, estimated around once a day. Nevertheless, these recourses are inherently unpredictable. A good relationship seems very important for an adequate care of the patients and a rational use of hospital resources. The collaboration between these two entities remains poorly described. Therefore, we aimed to characterize those recourses.

Patients and methods This is a prospective observational multicenter study conducted during 1 month (March 2016) in the ED and ICU of three hospitals of Paris suburb: Hôpital Antoine Béclère (398 beds university hospital), Hôpital André Mignot (702 beds) and Centre hospitalier intercommunal de Meulan les Mureaux (517 beds). We first conducted a preliminary survey describing the baseline evaluation of those recourses, with a questionnaire distributed to all physicians in the participating units. Thereafter, we did a prospective work to characterize the reality of these recourses. A questionnaire was fulfilled at each call, by the ED and the ICU physicians, blinded from his colleague. Finally, to evaluate the "non-recourse" too, we recorded the intensivist's evaluation. A questionnaire was distributed to all physicians involved in the participating units. Therefore, we did a prospective work to characterize the reality of these recourses. A questionnaire was fulfilled at each call, by the ED and the ICU physicians, blinded from his colleague. Finally, to evaluate the "non-recourse" too, we recorded the intensivist's evaluation. A questionnaire was distributed to all physicians involved in the participating units.

Results Preliminary survey: 40 emergency physicians and 24 intensivists answered (response rate: 87%). Intensivists declared that generally they estimate that the recourse is justified (87% of answers), in an appropriate timing (72%), and that they generally agree with the ED's physician about the emergency level (92%).

- Recourse characterization: during the study period, there were 111 recourses among the 12,674 patients admitted in the ED. We had bilateral data in 50% of cases. The mean recourse frequency was 1.2/day, representing 1/114 patients admitted in the ED, with a great variation between hospitals: from 1/71 to 1/158 patients. 50% of these recourses occurred during daytime (8 AM to 6 PM).

Intensivist went to the ED in 81% of cases. For the remaining cases, a direct admission was decided in 40% of cases and a phone advice was given in 60%. The mean time of arrival to the ED was 10 min and the mean time spent in the ED was 32 min. Life sustaining treatment limitation is a frequent reason for ICU admission (9.4%).

There was a non significant trend toward an estimation of higher severity by emergency physicians (37.7 vs 19.8%, p = 0.19). The exchange's climate was rated excellent or good in 81% of cases. It was more often "neutral" when an ED's resident was calling (45 vs 5.7% of recourses).

Discussion The ICU physicians seem to be solicited by the ED physicians roughly once a day. There is some "non-recourse" with delayed admission in ICU which could be deleterious for the patient. The climate of medical exchanges is usually good, although this appreciation may be downgraded by a resident's call.

Except for the Glasgow coma scale (which was suggested in our questionnaire), no severity or prognostic score was used as objective assessment of severity.

Life sustaining treatment limitation is a frequent reason for ICU recourse. These complex situations require a collegial discussion and recourses were estimated justified by the intensivists.

Conclusion Recourses to ICU physicians by ED physicians seem to be adapted and the medical exchanges happen in good relationship conditions. The frequency of these recourses is very variable among hospi- tal. ED physicians are mainly asking for an admission in ICU. Our work seems to point out some improvement ways, such as the use of severity scores, and a better supervision of the residents when asking for an intensivist's evaluation.

Competing interests None.

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O43 Intra hospital transport’s complications: incidence and risk factors

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Annals of Intensive Care 2017, 7(Suppl 1):O43

Introduction Caring for patients during intra hospital transport (IHT) is a high-risk activity. Adverse events during transport are frequent and may have significant consequences for the patient. The aim of this study was to assess the incidence of complications occurring during the IHT and to analyze the causes of such complications.

Patients and methods We prospectively describe IHT from the emergency department, realized from January 2016 to March 2016. Were included in the study IHT of compromised patients for whom critical care monitoring was needed and emergency physician is required. Clinical characteristics of patient’s departure and technical equipments (mechanical ventilation, drugs) were noted. Complications were defined as follows: patient related problems (desaturation, haemodynamic instability, arrhythmia, extubation, acute change in mental status, death) and ventilator related problems (breakdown or defect of the material).

Results During the inclusion period, 102 IHT were carried out. The IHT were realized for imaging procedure in 41 cases and for transferring patients to the intensive care unit in 24 cases and to the other wards in 37 cases. The median IHT duration was 15 min [10–30]. Twenty patients (19%) were mechanically ventilated. The majority of IHT (60%) were performed by the night shift emergency team. The incidence of complications was 44% (45 patients). Most events were related to haemodynamic instability in 25 cases, desaturation in 22 cases, agitation in 14 cases and cardiac arrest in 2 cases and one death. Therapeutic interventions were volume resuscitation in 13 cases, optimization of sedation in 12 cases, vasopressor management in 12 patients and cardiopulmonary resuscitation in 3 cases.

The occurrence of complications during transport was significantly increased in mechanically ventilated patients (p = 0.009), especially with inspiratory oxygen fraction >0.5 (p = 0.00), sedation before transport (p = 0.001), vasopressor requirement before transport (p = 0.03) and with the night shift team (p = 0.007). Sedation and mechanical ventilation were the independent risk factors of IHT complications.

Conclusion This study confirms that the intrahospital transport of compromised patients leads to a significant number of complications. This finding emphasises the need of improving medical skills during IHT.

Competing interests None.

O44 Influence of shift duration on cognitive performance of emergency practitioners: a prospective cross-sectional study

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Introduction Shift work including night work is responsible for sleep deprivation and tiredness. The relationship between tiredness and the risk of medical errors is now commonly accepted. The main objective of this study was to evaluate the cognitive performance of emergency practitioners after night shift of 14 h and after work shift of 24 h and to compare them with tests performed after a rest night at home.

Materials and methods We conducted an observational, prospective, single-center, randomized and cross-over comparative study. Emergency practitioners (staff physicians and residents) were eligible if they worked at least one night from 6:30 p.m. to 8:30 a.m. (14 h) and a day followed by consecutive night from 8:30 a.m. to 8:30 a.m. (24 h) within three consecutive months. Emergency practitioners participated to three cognitive assessments separated by at least 7 days: after a night of rest, after a night shift of 14 h and after a work shift of 24 h (including a night shift). The evaluation after night rest took place at least 3 days after the previous night shift. Three assessments were randomly assigned via sealed envelopes within a period of 3 months for each participant. Each participant was his own control and was evaluated by the same examiner. A psychologist formed four voluntary examiners to assess cognitive performance. Psycho-cognitive assessment began with self-evaluation of tiredness, attention, mood and lack of sleep (visual scales). Then an examiner evaluated participant’s cognitive performance according to the Wechsler Adult Intelligence Scale and the Wisconsin Card Sorting Test. Four cognitive skills were assessed: speed of processing information, working memory capacity, cognitive flexibility and perceptual reasoning. To test our main hypothesis, we performed an analysis of variance with repeated measures.

Results Forty emergency practitioners were included: 19 staff physicians and 21 residents. A staff physician and a resident declined to participate. Staff physicians were 36.2 ± 7.1 years old and residents were 26.8 ± 1.0 years old (p < 0.001). Average number of night shift per month was 4.4 ± 1.1 for staff physicians and 3.2 ± 0.6 for residents (p < 0.001). Amount of sleep, phone wakes up and stand ups were not different among staff physicians and residents. For all participants, no cognitive capacity was significantly altered after a night shift of 14 h when compared with performance after a night of rest. Conversely, three cognitive abilities were impaired after a work shift of 24 h when compared with performance after a night of rest: speed of processing information (11.2 ± 2.7 vs 12.4 ± 3.2; p < 0.003), working memory capacity (10.1 ± 2.9 vs 11.6 ± 3.0; p < 0.001) and perceptual reasoning (8.4 ± 2.7 vs 10.6 ± 2.8; p < 0.001). In absolute percentage, those performances were 10 to 20% lower after work shift of 24 h than after night of rest. Only cognitive flexibility scores were not significantly different in this condition. Working memory capacity was the only cognitive ability significantly altered after a work shift of 24 h versus a night shift of 14 h (10.1 ± 2.9 vs 11.6 ± 3.0; p < 0.01) whereas other cognitive abilities were all lower after a work shift of 24 h but without significant difference. Regarding the influence of medical experience, cognitive performance of staff physicians compared to residents were not statistically different after a night shift of 14 h and after a work shift of 24 h. There was no significant correlation between self and hetero assessment.

Conclusion The cognitive abilities of emergency practitioners were significantly altered following a work shift of 24 h whereas they were not significantly different from rest condition after a night shift of 14 h. Limiting 24 h shift work for emergency department should be considered and further evaluated.

Competing interests None.

O45 In hospital medical emergency calls: epidemiology and patients outcomes

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Introduction In hospital medical emergency (IME) is a high-risk activity. Adverse events during transport are frequent and may have significant consequences for the patient. The aim of this study was to evaluate the cognitive performance of emergency practitioners during transport (IHT) and to analyze the causes of such complications.

Materials and methods We conducted an observational, prospective, single-center, randomized and cross-over comparative study. Emergency practitioners (staff physicians and residents) were eligible if they worked at least one night from 6:30 p.m. to 8:30 a.m. (14 h) and a day followed by consecutive night from 8:30 a.m. to 8:30 a.m. (24 h) within three consecutive months. Emergency practitioners participated to three cognitive assessments separated by at least 7 days: after a night of rest, after a night shift of 14 h and after a work shift of 24 h (including a night shift). The evaluation after night rest took place at least 3 days after the previous night shift. Three assessments were randomly assigned via sealed envelopes within a period of 3 months for each participant. Each participant was his own control and was evaluated by the same examiner. A psychologist formed four voluntary examiners to assess cognitive performance. Psycho-cognitive assessment began with self-evaluation of tiredness, attention, mood and lack of sleep (visual scales). Then an examiner evaluated participant’s cognitive performance according to the Wechsler Adult Intelligence Scale and the Wisconsin Card Sorting Test. Four cognitive skills were assessed: speed of processing information, working memory capacity, cognitive flexibility and perceptual reasoning. To test our main hypothesis, we performed an analysis of variance with repeated measures.

Results Forty emergency practitioners were included: 19 staff physicians and 21 residents. A staff physician and a resident declined to participate. Staff physicians were 36.2 ± 7.1 years old and residents were 26.8 ± 1.0 years old (p < 0.001). Average number of night shift per month was 4.4 ± 1.1 for staff physicians and 3.2 ± 0.6 for residents (p < 0.001). Amount of sleep, phone wakes up and stand ups were not different among staff physicians and residents. For all participants, no cognitive capacity was significantly altered after a night shift of 14 h when compared with performance after a night of rest. Conversely, three cognitive abilities were impaired after a work shift of 24 h when compared with performance after a night of rest: speed of processing information (11.2 ± 2.7 vs 12.4 ± 3.2; p < 0.003), working memory capacity (10.1 ± 2.9 vs 11.6 ± 3.0; p < 0.001) and perceptual reasoning (8.4 ± 2.7 vs 10.6 ± 2.8; p < 0.001). In absolute percentage, those performances were 10 to 20% lower after work shift of 24 h than after night of rest. Only cognitive flexibility scores were not significantly different in this condition. Working memory capacity was the only cognitive ability significantly altered after a work shift of 24 h versus a night shift of 14 h (10.1 ± 2.9 vs 11.6 ± 3.0; p < 0.01) whereas other cognitive abilities were all lower after a work shift of 24 h but without significant difference. Regarding the influence of medical experience, cognitive performance of staff physicians compared to residents were not statistically different after a night shift of 14 h and after a work shift of 24 h. There was no significant correlation between self and hetero assessment.

Conclusion The cognitive abilities of emergency practitioners were significantly altered following a work shift of 24 h whereas they were not significantly different from rest condition after a night shift of 14 h. Limiting 24 h shift work for emergency department should be considered and further evaluated.

Competing interests None.
Introduction Medical emergency team is called to provide acute care for compromised patients outside intensive care unit. While waiting for his arrival, medical ward’s physicians must initiate patient rescue. The aim of this study was to describe clinical characteristics and outcomes of patients who experience medical emergency team calls and to evaluate initial care provided in medical specialty wards.

Patients and methods It’s a two-months prospective study (03/2016–04/2016) including all in hospital emergency calls. These calls were given by medical wards teams (dermatology, Gastroenterology, internal medicine, dermatology, cardiology and radiology) to the emergency physician when a deterioration in their patient’s condition was documented. Emergency and intensive care department emergency calls were excluded.

Results There were a total of 51 calls for 40 patients, principally hospitalized in internal medicine ward (21) and cardiology ward (14). Emergency calls were given by a physician in 47 cases and a nurse in 4 cases, during the night in 32 cases (62%). The greater number of calls was received Monday (13) and the week-end (11). Cardiac arrest (7), respiratory distress (28) and hemodynamic instability (16) were the reason’s call. The median emergency team arrival time was 10 min. The majority of patients (88%) required acute care for an average 35 min spent time. The vital signs were monitored and adequate therapy was initiated by the attending physician for 17 (53%) patients. Cardiovascular support was necessary for 24 (47%) patients, respiratory support was necessary for 16 (31%) patients. A transfer for the intensive care unit was indicated for 16 patients (31%) and delayed for 13 patients because of a lack of beds. In hospital mortality was 23% (n = 12). In hospital cardiac arrest and coma (Glasgow coma scale <8) were the principal prognostic factors. Delayed emergency time arrival was not associated with a greater mortality.

Conclusion In hospital emergencies were associated with an increased mortality. A better recognition of the instable patients may improve their prognosis.

Competing interests None.

O46 A new approach of sepsis heterogeneity Grégory Papin 1, Sébastien Bailly 2, Claire Dupuis 3, Stéphane Ruckly 4, Marc Gaurner 5, Laurent Argaud 6, Elie Azoulay 7, Adre Christophe 8, Bertrand Souweine 9, Dany Goldgran-Toledano 10, Guillaume Marcotte 11, Jean-Marie Forel 12, Romain Sonneville 13, Anne Sylvie Dumenu 14, Michael Darmon 15, Maïté Garroutte-Orges 1, Schwebel Carole 16, Jean-François Timst 17.

Background In the intensive care unit (ICU), the treatment and the outcome of patients with sepsis are heterogeneous. Identifying better homogeneous clusters of patients with sepsis and septic shock may improve their prognostic.

Aims We performed a national prospective multicenter ICU cohort. A first test set allowed the construction of more homogeneous clusters of patients with sepsis and septic shock. The second test set included 2017 patients admitted for sepsis or septic shock. We sought to identify new definitions of sepsis.

Methods We included 2017 patients admitted for sepsis in 12 French ICUs. We described the clinical characteristics of each cluster, 3 years mortality was compared with independent of patients’ prognosis. After description of maims characteristics of each cluster, 3 years mortality was compared with log rank test. A binary tree was built to assign new patients into cluster and evaluate using the validation set.

Results The test set included 4050 (67%) patients admitted for a first episode of sepsis. Six distinct clusters were identified (Fig. 7). Three years mortality was 21% (16–26%) for COPD exacerbation cluster, 26% (24–28%) for pulmonary sepsis cluster, 37% (33–40%) for surgical sepsis cluster, 27% (19–34%) for meningoco-encephalitis cluster, 47% (41–52%) for immunocompromised patients cluster and 47% (44–50%) for chronic diseases cluster (p < .001). Binary tree identified 6 discriminant variables to assign patients into clusters: lung infection, surgical admission, bronchial infection, meningone infection, hematological malignancy and chronic heart failure. Identical patient profiles were found in the validation set.

Conclusion Six clusters of ICU patients admitted for sepsis or sepsis shock were identified. All clusters contained patients who met the new definitions of sepsis. Despite this, these clusters have a very high heterogeneity prognosis. Considering these clusters may improve homogeneity of patient’s enrolled in future clinical trials.

Competing interests None.

O47 Antimicrobial strategy for severe community-acquired Legionnaires’ disease: a multicentre retrospective observational study Jerôme Cecchini 1, Samuel Tuffet 2, Romain Sonneville 3, Muriel Fartoukh 4, Julien Mayix 5, Damien Roux 6, Amandine Koutouchi 6, Florence Boissier 6, Martial Chui 5, Martial Thyraud 6, Eric Maury 6, Sébastien Jochmans 1, Mékontso Dessap Armard 2, Christian Brun-Buisson 1, Nicolas de Prost 1.

Background Legionnaires’ disease (LD) is a pneumonia caused by Legionella pneumophila and remains a severe illness. The optimal management of LD remains controversial.

Aims To compare the outcome of patients according to the antibiotic strategy administered: (1) fluoroquinolones versus no fluoroquinolones and (2) monotherapy versus combination therapy.

Methods In a multicenter study including patients admitted to 10 ICUs over 1 year period, 124 patients were included. Patients were stratified according to the antibiotic therapy administered: (1) fluoroquinolones versus no fluoroquinolones and (2) monotherapy versus combination therapy.

Results Of all patients, 61% were treated with fluoroquinolones and 54% had a monotherapy. The outcome was not associated with the antibiotic strategy. No significant difference was found in terms of mortality, but patients treated with fluoroquinolones had a higher incidence of acute kidney injury.

Conclusion Our study suggests that fluoroquinolone monotherapy is associated with a higher incidence of acute kidney injury. Further studies are needed to evaluate the optimal treatment of LD.
Table 7 Univariable and multivariable Cox model of factors associated with day-60 mortality in patients (n = 211) with severe Legionnaires’ disease admitted in the ICU

| Combination therapy | Univariable analysis |
|---------------------|----------------------|
|                     | HR (95% CI), p value  |
| No                  | 1                    |
| Yes                 | 0.56 [0.32–0.97], p = 0.04 |

Fluoroquinolone therapy

|                        | HR (95% CI), p value |
|------------------------|----------------------|
| No                     | 1                    |
| Yes                    | 0.38 [0.21–0.68], p = 0.001 |

Age > 60 years

|                        | HR (95% CI), p value |
|------------------------|----------------------|
| No                     | 1                    |
| Yes                    | 2.74 [1.37–5.50], p = 0.004 |

Smoking

|                        | HR (95% CI), p value |
|------------------------|----------------------|
| No                     | 1                    |
| Yes                    | 0.33 [0.17–0.64], p = 0.001 |

LODS (per point)

|                        | HR (95% CI), p value |
|------------------------|----------------------|
| No                     | 1                    |
| Yes                    | 1.34 [1.23–1.46], p < 0.0001 |

Propensity score (per point)

|                        | HR (95% CI), p value |
|------------------------|----------------------|
| No                     | 1                    |
| Yes                    | 0.23 [0.09–0.57], p = 0.002 |

I/NR included not retained, LODS Logistic Organ Dysfunction Score, HR hazard ratio

* Propensity score of fluoroquinolone-based treatment

combination therapy. To evaluate the effect of antimicrobial strategy on ICU mortality, multivariable Cox model and propensity score analyses were used.

Results 211 patients with LD requiring ICU admission were included. A fluoroquinolone-based and a combination therapy were administered to 159 and 123 patients, respectively. 146 patients (69%) developed an acute respiratory distress syndrome, 111 (53%) a shock, 56 (27%) an acute renal failure that required renal replacement therapy, and 54 (26%) died in the ICU. In-ICU mortality was lower in the fluoroquinolone group as compared to the no-fluoroquinolone group (21 vs. 39%, p = 0.01), as well as in the combination therapy group as compared to the monotherapy group (20 vs. 34%, p = 0.02). In multivariable analysis with covariate adjustment including a propensity score for fluoroquinolone treatment, a fluoroquinolone-based therapy, but not a combination therapy, was independently associated with a reduced risk of mortality (HR 0.4, 95% CI [0.2–0.7], p = 0.002) (Table 7).

Discussion Our results, which suggest a beneficial effect on mortality of a fluoroquinolone-based therapy in patients with LD who required ICU admission, are consistent with previous studies showing a non-significant trend in favor of fluoroquinolone use in patients hospitalized for LD [1]. The limited number of patients in our study precluded assessing the individual effects of distinct molecules within the fluoroquinolone class.

Conclusion Patients with severe LD receiving a fluoroquinolone-based antimicrobial regimen in the early course of the management in ICU had a lower in-ICU mortality, which persisted after adjusting on significant covariates. A combination therapy did not provide significant mortality benefit in the current study.

Competing interests None.

Reference
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O48 Necrotizing pneumonia in ICU: clinical, microbiological and radiological characteristics

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Introduction Data on necrotizing pneumonia in ICU are scarce. This potentially devastating condition is generally thought to be due to highly virulent bacteria such as Panton-Valentine leukocidin-secreting Staphylococcus aureus or to slowly-growing organisms such as mycobacteria and other Actinomycetales. The goal of this study was to describe the clinical, microbiological and radiological features of NP in intensive care unit.

Patients and methods Monocenter retrospective study. All patients hospitalized between January 1st 2009 and July 1st 2016 with an ICD-10-CM code for pneumonia (U11) in an 18-bed medical ICU were included. All chest X-rays and CT-scans obtained during the stay were screened in order to select patients with NP, which was defined as a new cavity or a zone with no contrast enhancement within a pulmonary consolidation. Diagnosis was confirmed by a senior radiologist. Medical records of included patients were reviewed to describe clinical and radiological presentation, demographic characteristics, microbiological results and outcome.

Results Among 1009 screened pts, a definitive diagnosis of NP was made in 37 pts (3.7%). Median age was 57 [interquartile range 48–69] years, 68% were males, median SAPS2 was 43. Half of the pts were immunocompromised due to hematologic malignancy (7 pts), solid tumor (7 pts), immunosuppressive therapy (2 pts) or HIV infection (2 pts). Eight pts had chronic respiratory disease, 8 had diabetes, 11 were alcoholic.

Pneumonia was community-acquired in 21 pts, including 3 pts with aspiration pneumonia. The remaining NP were nosocomial diseases, including 5 pts with ventilator-associated pneumonia. In 4 pts, pneumonia was considered to be acquired by hematogenous spread, including 1 patient with endocarditis.

Most pts had severe hypoxemia, with a median minimum PaO2/FIO2 ratio of 153 [102–203] mmHg. Twenty-three pts were intubated, and median duration of mechanical ventilation was 4 days. Eighteen pts developed septic shock, and 3 had renal replacement therapy. In-ICU mortality was 32% (12 pts).

Chest CT-scan was performed in all pts but 1. One or more cavity was observed in 33 pts, whereas lung necrosis without excavation was present in 4 pts. Most pts (25) had more than one lesion, involving more than 1 pulmonary lobe in 22 pts and both lungs in 15 pts. Median size of the largest lesion was 42 [24–78] mm.

Broncho-alveolar lavage was performed in 25 pts, and bronchial aspirate in 18 pts. One or more pathogen was isolated in 29 pts; in 4 pts, the final diagnosis was non-infectious (neoplasia in 3 pts and 1 crack lung), and all samples were sterile in 4 additional pts. Fifteen pts had polymicrobial infection. Overall, the most frequently isolated pathogen was Pseudomonas aeruginosa (9 pts), followed by Staphylococcus aureus (7 pts) and fungi (7 pts). Klebsiella pneumoniae and other Enterobacteriaceae were isolated in 5 and 6 pts, respectively. All cases involving P. aeruginosa were diagnosed in immunocompromised patients. In contrast, most patients with S. aureus infection (6/7) and all patients with K. pneumoniae infection were immunocompetent.
Among pts with proven infection, none underwent surgical treatment, but percutaneous drainage was performed in 3 pts.

**Conclusion** NP was rarely observed in our ICU. In immunocompromised patients, *P. aeruginosa* and fungi were the most frequently isolated pathogens, whereas highly virulent bacteria such as *S. aureus* and *K. pneumoniae* were more frequently encountered in immunocompetent patients.

**Competing interests**

None.

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**O49 Epidemiology and long-term outcome after severe symmetric peripheral gangrene**

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**Annals of Intensive Care 2017, 7(Suppl 1): O49**

**Introduction** Symmetric peripheral gangrene (SPG) is a rare but severe complication of septic shock often leading to multiple amputations. Epidemiology of SPG and long-term outcome remain poorly known. Amputations are serious offense to body integrity but can benefit from rehabilitation and prosthetics. Our objectives were to describe epidemiology of SPG and to assess health-related quality of life (HRQOL) once rehabilitation was achieved.

**Patients and methods** A prospective and retrospective, multicentric study was performed. Adult patients hospitalised between 2005 and 2015 were included. They must have undergone at least two major amputations (whatever the level) and been discharged in a specialised rehabilitation center. HRQOL was assessed with generic scale EQ-5D-3L by phone call. Epidemiologic data were extracted from hospitalisation reports.

**Results** Nine centres on 13 participated, 30 patients were recruited and medical letters were available for 25 of them. SPR was observed in a majority of female (60%), aged around 50 yo. Mean intensive care unit (ICU) length of stay was 39 (± 21) days. Infectious agents were in majority Gram positive cocci (64%), but Escherichia Coli took an important part (20%), similar to Meningococcus (16%). All patients were amputated of the two lower limbs and 80% were quadruple amputees. HRQOL estimated wit EQ index was inferior to the French reference. However, patients rated themselves their health state as similar to the reference and even superior to the reference French value before SPR, using visual analogue scale (VAS). Main decrease in VAS was 22 points (95% CI 13–31). Intense pain due to phantom pain was the main factor of impaired EQ index. Painkillers use was statistically dependant of antidepressants use. All patient except one said they would be willing to be treated again for SPG.

**Conclusion** Symmetric peripheral gangrene is mainly due to Gram positive Cocci but also Escherichia Coli. It leads to severe amputations with impaired HRQOL which could be improved by better analgésic strategies. However patients consider themselves as being in good health and would be willing to be treated again. This should be taken into account before withdrawing life-sustaining therapies.

**Competing interests**

None.
time of antimicrobial therapy for reduction of bacterial inoculum. However, valve surgery in patients with positive valve culture was not associated with death or post-surgery complications except ARDS and clinicians must be aware of this risk.

**Conclusion** Our results suggest that valve cultures are more frequently positive when surgery is performed before 7 days of adequate antimicrobial therapy. We did not find that positive valve culture was associated with worst ICU outcomes except for ARDS.

**Competing interests** None.

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**O51 Time course of septic shock in immunocompromised and non-immunocompromised patients**

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**Annals of Intensive Care 2017, 7(Suppl 1):O51**

**Introduction** The outcome of septic shock has undoubtedly improved over the last two decades, both in immunocompetent and immunocompromised patients. The combination of anti-infective treatments and aggressive organ failure supports often allows stabilisation of the clinical condition, but patients become then exposed to intensive care unit (ICU)-acquired infectious and non-infectious complications that significantly impact on prognosis. Whereas immunosuppression is commonly viewed as a major risk factor of death in septic shock, whether it is associated with an increased incidence of ICU-acquired complications is unclear. The present study aimed at addressing the course of septic shock across clinically significant subgroups of patients with and without immunosuppressive conditions.

**Patients and methods** This was a 8-year (2008–2015) monocenter retrospective study performed in a 24-bed medical ICU. All consecutive adult patients diagnosed for septic shock within the first 48 h of intensive care unit (ICU) admission were included. Septic shock was defined as a microbiologically proven or clinically suspected infection, associated with acute circulatory failure requiring vasopressors despite adequate fluid filling. Patients were considered either as non-immunocompromised or immunocompromised, this latter subgroup being categorized in three subgroups according to the underlying immunosuppressive condition: solid neoplasm, hematological malignancy and HIV- or drug-induced cellular immunosuppression. Furthermore, administration of intravenous chemotherapy during the last 3 months and leucopenia defined by leucocyte count <1000/mm³ and/or neutrophil count <500/mm³ were collected as well. Survival status was assessed at ICU and hospital discharge. We focused on the most likely infectious, hemorrhagic and ischemic complications occurring after the first 48 h in the ICU. The impact of the underlying immune status on 3-day and late mortality was assessed using a cause-specific proportional hazard model. To assess the impact of immune status on ICU complications, we performed two competing risk analysis model (Fine-Gray and cause-specific proportional hazard model). A landmark analysis with the choice of a 3-day fixed landmark point was performed to correct for the substantial immortal time bias. The SAPS2 was used to adjust for admission severity.

**Results** Eight hundred one patients were included. Their median age was 59 (57–79) years. The main source of infection was the lung in 49.6%. About 38% of patients had underlying immunosuppressive conditions, distributed into solid neoplasms (15.2%), hematological malignancy (13.2%) and non malignant immunosuppression (9.6%). The overall in-ICU and in-hospital mortality rates were 37.3% and 41.3%, respectively. With respect to the immune status, immunocompromised patients displayed worse outcome than immunocompetent ones, those with solid neoplasms having the highest risk of death. 113 patients died within the first 3 days, leaving 633 patients at risk of ICU-acquired complications. Among the 3-day survivors, the crude incidence rates of ICU-acquired infectious, ischemic and hemorrhagic complications were 27, 11 and 9%, respectively. The incidence of secondary infections was similar across the immunocompetent and the three immunocompromised subgroups of patients. In contrast, non malignant immunosuppression was associated with an increased risk of ischemic complications while hematological malignancies were associated with an increased risk of bleeding.

**Conclusion** We herein show that patients with septic shock display various clinical courses in relation with their underlying immune status. The burden of ICU-acquired complications in septic shock calls for every efforts of prevention and early detection, in order to improve the overall outcome.

**Competing interests** None.
[n = 47 (87%),] oral mucositis [n = 36 (66.7%),] paralytic ileus [n = 17 (31.5%),] nausea and vomiting [n = 14 (25.9%),] Digestive hemorrhage occurred in 9 (16.7%) patients. Almost all patient had abdominal CT-scan (94.4%). The digestive inflammation involved ascending colon [n = 41 (75.9%),] transverse colon [n = 25 (46.3%),] descending colon or sigmoid [n = 45 (46.3%),] and small intestine [n = 12 (22.2%).] A diffuse involvement (more than one segment) occurred in n = 32 (59.3%) patients. The other frequent signs were a peritoneal effusion n = 37 (68.5%) and intra-abdominal fat infiltration n = 19 (35.2%). Pneumoperitonea (n = 2) and parietal panniculitis (n = 1) were rare.

Antibiotic therapy consisted in beta lactam (carbapenems [n = 31 (57.4%),] ureidopenicillins and inhibitors [n = 18 (33.3%),] cefepime [n = 4 (7.4%),] ceftazidime [n = 1 (1.9%)]) combined with aminosides [n = 41 (75.9%),] vancomycin [n = 44 (81.5%)], and ornidazole [n = 34 (63%)]. Antifungal therapy was used for 33 (61.1%) patients. There was 33 (61%) bacterial and 8 (14.8%) fungal documented infections. Empirical antimicrobial therapy was effective in respectively 31 and 3 cases. Most frequent pathogens were Escherichia coli (n = 10), Enterococcus faecium n = 8, Enterobacter cloacae (n = 7), Klebsiella pneumoniae (n = 6), Pseudomonas aeruginosa (n = 3), Candida spp (n = 5). Infection was polymicrobial for n = 16 (29.6%). Involvement of small intestine including terminal ileum was associated with fungal infections (p = 0.01). ICU mortality was associated with fungal infections (p = 0.003) and polymicrobial infections (p = 0.03). Five patients underwent surgery (2 perforations, 1 digestive hemorrhage, 1 abscess, 1 negative laparotomy), with a ICU mortality of 60%.

Conclusion NE is a life threatening complication in patients with malignancies who receive intensive chemotherapy. NE is associated with high mortality rates, especially in patients with fungal infections. Antifungal therapy should be systematically discussed in NE patients with small intestine involvement.

Competing interests None.

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O53 Early infectious complications following heart transplantation Stéphanie Pons1, Romain Sonneville2, Lenka Styfola2, Lila Boudama1, Mathilde Neuvillé1, Eric Maricotte1, Agnès Radjou1, Jordane Lebut1, Bruno Mourvillier1, Richard Dorent1, Marie-Pierre Dilly5, Patrick Nataf6, Michel Wolff1, Jean-François Timis1 1Réanimation médicale et infectieuse, Hôpital Bichat-Claude Bernard (AP-HP), Paris, France; 2ICuresearch, ICUREsearch, Paris, France; 3Cardiologie, Hôpital Bichat-Claude Bernard (AP-HP), Paris, France; 4Anesthésie réanimation chirurgie cardio-vasculaire, Hôpital Bichat-Claude Bernard (AP-HP), Paris, France; 5Chirurgie cardiovasculaire, Hôpital Bichat-Claude Bernard (AP-HP), Paris, France

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Introduction Heart transplantation is the reference treatment of end-stage heart failure. National priority heart transplantation for severe acute heart failure has increased, leading to a change in the population of heart-transplant recipients (older patients with more comorbidities). Infectious complication is one of the biggest concerns after solid organ transplantation, with high rates of morbidity and mortality. The impact of early infectious complications has never been assessed in a large cohort of ICU patients. We aimed to determine the characteristics, the determinants and the impact of infectious complications following heart transplantation.

Patients and methods We retrospectively studied all consecutive heart-transplant recipients from Bichat university hospital, between January 1st, 2011 and June 30th, 2015. All infectious complications that occurred within 6 months after transplantation were considered for analysis. The early post-operative period was defined as the first 6 days following heart transplantation. The primary endpoint was the rate of infectious complications at 6 months. We used multivariate logistic regression to identify independent risk factors for infections at 3 months was determined by Cox regression analysis. Data are presented as median (interquartile range) or number (percentage).

Results One hundred and thirteen patients (53 years [40–62], male n = 62 (76.1%),) were included. At the time of heart transplantation, 65 (57.5%) patients were hospitalized in ICU for severe acute heart failure and 28 (24.8%) patients were under extracorporeal membrane oxygenation (ECMO) support. The SOFA score on the day before heart transplantation was 4 [1–6]. A cytomegalovirus (CMV) mismatch was found in 15 (14%) recipients. Twenty-two (19%) patients had a multidrug-resistant bacteria (MRB) carriage before transplantation, and six (5.3%) acquired one during the first week in the ICU. There were 213 infectious complications within 6 months after heart transplantation in 90 (80.5%) patients. The first bacterial infection occurred in ICU for 66 (58.4%) patients. Pneumonia was the most frequent infection, accounting for 46.5% of the cases. Fifty-six pneumonia were diagnosed during the ICU stay, and 46 out of 65 (70%) within the early post-operative period. By contrast, bloodstream infections (n = 31, 14.5%) and Scarpas infections (n = 22, 10.3%) were as frequent in the early post-operative period than after day eight. Sixty-three patients (55.8%) developed at least one infection due to Gram-negative bacilli, including 24 patients (21.2%) with an infection due to Pseudomonas aeruginosa. Thirty-five patients (31%) had a Gram-positive cocci infection. Multi-resistant bacteria were responsible for infection in twenty-one (16.8%) patients. Within 6 months following heart transplantation, 44 (38.9%) patients had a viral complication, usually occurring after the early post-operative period and before day 30 (n = 32 [28.3%]). Fungal infection was found in 16 (14.2%) patients, including seven invasive aspergillosis, most of them diagnosed after day 30. In univariate analysis, we found that ECMO following heart transplantation (n = 10 [29.4%] vs n = 62 [49%] p < 0.01), day 1 SOFA score (8 [7–10] vs 9.5 [8–11] p = 0.01) and mechanical ventilation duration (3 [1–8] vs 9 [5–15] p < 0.01) were associated with infectious complications in ICU. After adjustment, ECMO following heart transplantation was identified as the only independent risk factor for bacterial infection in the ICU (Odds Ratio: 3.1, 95% confidence interval 1.4–6.9, p = 0.006). Bacterial infection in ICU was associated with a longer stay in ICU (20 [13–29] days versus 12 [8–17] days, p < 0.001), but not with ICU mortality. Ninety-two patients (81.4%) were alive 6 months after the transplantation, and 11 out of the 21 deaths (52.3%) were caused by an infectious complication. In an adjusted Cox model, the third not-viral infection was significantly associated with death at 3 months post transplantation (adjusted Hazard Ratio: 6.2, 95% confidence interval 1.2–31, p = 0.02).

Conclusion This study confirms the high rate of early infectious complications after heart transplantation. ECMO following transplantation is an independent factor associated with bacterial infections. Bacterial infection in ICU was associated with a longer stay in ICU. The risk of death at 3 months after transplantation increased dramatically with the third episode of “not-viral infection”.

Competing interests None.

O54 Clinical course and prognosis of severe cryptococcosis in the intensive care unit: a retrospective multicenter study Aëlle Le Gall1, Mathilde Neuvillé1, Simon Bouvier1, Julien Mayaux2, Damien Contou3, Yacine Tandjaoui-Lambiotte4, Vincent Das5, Benjamin Zuber6, Stéphane Gaudry1, Guillaume Voisiot7, Mikhail Alves1, Eric Maury1, Naike Bigi1 1Réanimation médicale, Hôpital Saint-Antoine, Paris, France; 2Service de réanimation médicale et infectieuse, Hôpital Bichat-Claude Bernard-APHP, Paris, France; 3Réanimation médicale, Hôpital Pitié-Salpêtrière, Paris, France; 4Réanimation Médicale, Hôpital Henri Mondor, Créteil, France; 593, Hôpital Avicenne, Bobigny, France, 6Réanimation polyvalente adulte, Centre Hospitalier Intercommunal André Grégoire, Montreuil, France; 7Intensive care unit, Hospital Center De Versailles, Le Chesnay, France; 8Intensive care unit médico-chirugirale, CHU Louis Mourier, Colombes, Colombes, France;
Competing interests

Introduction Cryptococcosis is a well-known opportunistic infection, especially in HIV-infected patients with profound immunosuppression. However, data regarding presentation and prognosis of severe forms of cryptococcosis requiring intensive care unit (ICU) admission are scarce.

Patients and methods We performed a retrospective multicenter study including all patients admitted for cryptococcosis from January 1998 to March 2016 to one of the 11 participating ICUs. Admissions were identified through a systematic review of ICUs databases using ICD-10 code B45. Diagnosis was confirmed either by cryptococcal antigen detection in serum, cerebrospinal fluid, bronchoalveolar lavage or urine, or by the identification of Cryptococcus in the culture of a specimen of any site or by histology. Qualitative and quantitative values are expressed as number and percentage, and median and interquartile range, respectively. Comparisons between ICU-survivors and non-survivors were performed using Fisher’s exact test and Mann-Whitney test for qualitative and quantitative variables, respectively. A p-value < 0.05 was considered to be significant.

Results Seventy-three patients (age 44 [37; 52] years, 58 [79.5%] male, SAPS II: 42 [31; 59], SOFA score at day one: 4 [1;7]) were included. Twenty-five (34.2%) of them reported previous opportunistic infection. HIV infection was the leading cause of immunosuppression (n = 58 [79.5%], median CD4 count: 12 [6; 40/mm3]) and was newly diagnosed in 30 (51.7%) patients. The remaining patients broke as follows: immunosuppressive therapy for auto-immune disease (n = 6) or solid organ transplantation (n = 4) or nephrotic syndrome (n = 1), cirrhosis (n = 1), primary hypogammaglobulinemia (n = 2) and diabetes mellitus as the only underlying comorbidity (n = 1). Impaired consciousness (68.5%) was the first reason for ICU admission, followed by acute respiratory failure (21.9%), shock (8.2%) and acute renal failure (1.4%). Time from first symptoms to diagnosis was 7 [3; 21] days, and was 9 [4; 25] days from first symptoms to ICU admission.

Central nervous system was the most frequently infected site (n = 62 [84.9%]) with meningoc-encephalitis (n = 60 patients) and brain abscesses (n = 8). Lung was the second more frequent site of infection (n = 14). Other localisations included skin, heart, urinary tract, pleura and spleen. In 23 patients (31.5%), at least two sites were infected. Fungemia was observed in 17 patients (23.2%). Invasive mechanical ventilation was required in 34 (46.6%) patients, vasopressors in 18 (24.7%) and renal replacement therapy in 10 (13.7%). Median ICU length of stay was 4 [3; 12] days. ICU mortality was 41.1%. Factors associated with ICU death were: infection of at least 2 sites (OR 5.1 [1.8; 15.2], p = 0.0038), SAPS II and SOFA score at day one (p < 0.001), need for invasive mechanical ventilation (OR 18.9 [5.6;63.3], p < 0.001), need for vasopressors (OR 41.3 [10.3;769.6], p < 0.0001) and renal replacement therapy (OR 7.5 [1.5;38.2], p = 0.013). Neither HIV status (p = 0.77) nor previous opportunistic infection (p = 0.80) was associated with ICU mortality.

Discussion Because of the limited number of non-HIV-infected patients, the study was not powerful enough to analyze clinical presentation according to HIV status.

Conclusion Severe forms of cryptococcosis requiring ICU admission are mostly observed among HIV-infected patients. However, one in five patients has another cause of immunosuppression, principally immunosuppressive therapy for auto-immune disease or solid organ transplantation. Central nervous system is the most common site infected. ICU mortality is high and associated with disseminated disease and need for organ supports. HIV status does not influence prognosis.

Competing interests None.
of this study is to determine recent national trends in occurrence of septic shock (SC) and SC-related deaths and costs.

**Patients and methods** We analysed the occurrence of sepsis from 2009 to 2014, in adult patients, using the national French hospital Database PMSI (Programme de Médicalisation des Systèmes d’Information). This database collects annually, for each hospital stay in France, a uniform hospital discharge record including diagnostic codes according to the International Classification of diseases, Tenth revision, Clinical Modification (ICD-10th-CM) and procedure codes according to the CCAM (Classification Commune des Actes Médicaux). Cases were identified from discharge records in the PMSI database. Septic shock was defined by a combination of a principal or accompanying diagnosis of infection (ICD-10thCM) plus a diagnosis code for septic shock (R572, R578, R579) or the procedure code EQLAFF03 (infection of vasopressors). Charlson index were determined using diagnosis codes as previously described [1].

Costs were determined according to the National Reference Costs (ENC = Etude Nationale des Couts). Temporal trend were assessed by calculating the value of the entire variable by year and compared using linear regression variables or Cochran Armitage Trend Test for continuous and categorical variables respectively.

**Results** A total of 25,444,627 adults aged ≥ 18 years were hospitalized in France during 2009–2014. Among them we identified 419,597 (10.3%) patients with SC. There was an annualized increase in the incidence of SC from 87 to 122/100,000 inhabitants p < 0.001. Only 25% of the cases were hospitalized in an affiliated university centre. Sepsis was more frequent among men (62.8%) than among women (37.2%). Overall the SAPS2 score was (median [interquartile range = IQR] 49 [36–65]. Charlson comorbidities index decreased significantly over time (p < 0.001). 58.6% of the cases were considered as medical. Nearly all the patients were admitted from the emergency department (75.9%) or from wards (20.1%). About 80% of the patients were hospitalized at least 1 day in Intensive Care Units and of note this percentage decreased consistently over time from 85.6 to 79.7%. Concomitantly admission at least 1 day in intermediate care unit (from 24.9% to 33.6%) or in specialized wards such as oncology-hematology (from 0.6% to 3.1%) increased over time. Overall, the main comorbidities reported were congestive heart failure (27.6%), cerebro-vascular disease (8.7%), Chronic Pulmonary Disease (15%), dementia (2.9%), renal disease (11.3%), diabetes without (6.3%) and with complications (5%) and cancer (16.9%), mild liver disease (8.5%). The main sources of sepsis were pulmonary (39.2%), abdominal (12%), urinary (15.6%), and cutaneous (3.2%), cardiac (3.1%), neurological (1.1%), and oesto-articular (1.6%). Gram-negative bacteria were the predominant organisms causing sepsis (from 27.6% to 38.8%) but gram-positive bacteria were reported most commonly in each subsequent year as well (from 20.8% to 26.4%). The length of ICU and hospital stays were 9 [3–19] and 18 [8–35] days respectively and both decreased over time (p < 0.001). Mortality rates slightly decreased over the 6-year period (from 41.7% to 38.7%), However the in-hospital deaths related to sepsis increased over time from 433 to 571/10,000 deaths in France (p < 0.001). The average costs per case were 14,453 [7460–25,041] decreasing over time from 15,291 [7637–26,136] to 14,171 [7593–24,194].

**Conclusion** SC incidence increased in France. Nearly 75% of the cases were managed in non-affiliated university centre. Although the overall mortality rates in patients with SC is slightly decreasing in France, the number of deaths related to SC in France is growing. Results must be interpreted with caution because of the limits related to administrative databases. However, the trends are comparable to data from other countries.

**Competing interests** None.

**Reference**

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**O57**

**Severe complications of Zika virus infection during the 2016 outbreak in Guadeloupe**

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**Introduction** Since 2015, Zika virus (ZIKV) infection is spreading across South and Central America. The outbreak reached the Caribbean in the early 2016. The aim of this study is to describe the characteristics and outcome of patients with severe complications of ZIKV requiring intensive care unit (ICU) admission.

**Patients and methods** We performed a prospective observational study in the ICU in the University Hospital of Guadeloupe from May 15th to September 15th 2016. All patients with an acute ZIKV infection confirmed either by Real Time PCR (RT-PCR) or IgM detection in the serum were included. In patients presenting with an acute neurologic disease, extensive laboratory testing, CSF analysis, brain MRI and electromyography testing were performed according to a predefined protocol, in order to rule out others diagnosis. Patients with Guillain-Barré syndrome (GBS) were treated with intravenous immunoglobulins 0.4 mg/kg/day during 5 days.

**Qualitative data are presented in percentage and quantitative data in median [interquartile range].**

**Results** During the study period, 19 patients with proven ZIKV infection were admitted to the ICU: 10 (53%) men, median age 59 [37–73] years, SAPS-2 30 [21–37]. Main comorbidities were chronic hypertension (47%) and diabetes (32%). One patient (5%) was immunocompromised and 6 (32%) had no comorbidities.

Among the 19 patients, 15 (79%) were admitted for an acute neurological disease, namely GBS in 10 (53%) patients and encephalitis in 5 (26%) patients. The remaining 4 (21%) patients were admitted for COPD exacerbation, lupus exacerbation, myasthenia gravis and metformin intoxication complicated with auto-immune hepatitis. Clinical symptoms of ZIKV infection such as fever, arthralgia, cutaneous rash, conjunctivitis were identified in 11 (58%) patients. The time between symptoms onset and ICU admission was 7 [5–10] days. Among the 15 patients with neurological diseases, the time between the onset of neurological symptoms and ICU admission was 3 [2–5] days. ZIKV RT-PCR was positive in urine in 15 (79%) patients, in blood in 2 (11%) patients and in CSF in 1 (5%) patient. Serum IgM were detected in 10 (53%) patients.

CSF was obtained in the 15 patients with neurological diseases. For the 10 patients with GBS, white cell count was 3 [1–5]/mm³; protein level was 1.4 [0.8–1.9] g/L; glucose level 4.8 [4.3–5.3] mmol/L. For the 5 patients with encephalitis, white cell count was 65 [24–180]/mm³; protein level was 0.6 [0.4–1.1] g/L, glucose level 5.5 [4.1–6.1] mmol/L. Brain MRI abnormalities were observed in 4 (21%) patients, and meningitis was observed in one patient.

Fourteen patients (79%) had mechanically ventilation for median length of 12 [6–22] days. Among them 4 (21%) were tracheotomised because of prolonged mechanical ventilation. Six (32%) patients received vaso-active drugs, and 2 (11%) required renal replacement therapy. ICU length of stay was 12 [7–25] days. Five (26%) patients stayed more than 30 days in the ICU: 3 patients with GBS, 1 patient with encephalitis, 1 other patient. Three (16%) patients died in the ICU.

**Conclusion** As previously reported, ZIKV infection is responsible for neurological complications [1]. In particular, the GBS has been described previously in a similar setting [2]. Our data confirmed the association between ZIKV infection and GBS, as the annual incidence of GBS in our ICU is usually 3–5. Interestingly, several cases of encephalitis were observed, which has not been reported previously.
These are preliminary data. As the outbreak is still ongoing in the Caribbean, we can expect more cases and more precise data within the next months.

**Competing interests**

None.

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O59

Closed-loop adjustment of oxygen flow with FreeO2 in patients with acute coronary syndrome, comparison of two SpO2 target and manual adjustment: a randomized controlled study

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**Introduction**

Supplemental oxygen has been used in the management of patients with acute coronary disease and investigated for more than a century [1]. The rational for a systematic oxygen supply in patients with acute coronary syndrome is based on a limited number of data. The links between hypoxia and ischemic or arrhythmic electrocardiographic abnormalities have been known for a long time. The risks associated with hyperoxia have been well described in physiological studies (increased coronary resistances, decreased coronary blood flow). A recent large RCT demonstrated that liberal oxygen administration during the acute phase of myocardial infarction may increase the level of cardiac enzyme and the infarct size [2].

FreeO2 is a recently developed device that automatically titrates oxygen flow with the aim to maintain constant the SpO2, around the target set by the physician. We evaluated two SpO2 targets using automated oxygen titration with FreeO2 during acute coronary syndrome and evaluated oxygenation delivered with usual manual oxygen therapy. Our hypotheses were that high rates of hypoxemia and hyperoxia would occur with manual titration, and would be reduced with automated oxygen titration.

**Patients and methods**

We conducted a pilot randomized controlled, single blind monocentric study to evaluate oxygen therapy administration at the acute phase of acute coronary syndrome. Patients with acute coronary syndrome were included (based on AHA criteria). Severe COPD patients were excluded. Patients were randomized one of the three arms: Automated oxygen titration with FreeO2 at two different SpO2 targets (92% and 96%) and manual administration of oxygen.

The study lasted a maximum of 48 h for each patient, including one night. All patients were continuously monitored with FreeO2, (set in recording mode in the manual group), continuous cardiac telemetry was performed for all patients, cardiac enzymes were collected as per usual care.

The primary outcome was the frequency of desaturation (SpO2 < 90% for 30 s). The secondary outcomes were the frequency of arrhythmias, the rate of tachycardia episodes and the level of cardiac enzymes in patients with acute coronary disease.

**Results**

Sixty patients were included in the study, the mean age was 63 ± 12 years, 73% of the patients were men. The average recording time was 11.5 ± 2.8 h. Preliminary data are presented here.

−20 patients were included in the FreeO2 group with SpO2 target = 92%.

−19 patients were included in the FreeO2 group with SpO2 target = 97% (including one case of missing data due to technical issues).

Primary end point (rate of hypoxemia): The percentage of time with hypoxemia (SpO2 < 90%) was 4% of the recording time (equivalent to 30 min) in the control group, 1.2% of the recording time (equivalent to 9 min) in the FreeO2 (92%) group and 0.4% of the recording time (equivalent to 2.5 min) in the FreeO2 (97%) group (manual vs. 92%, p = 0.58; manual vs. 97%, p = 0.001; 92 vs. 97%, p = 0.006).

57% of the patients in the control group, 30% in the FreeO2 (92%) and 22% in the FreeO2 (97%) experienced desaturation below 90% for more than 30 s (manual vs. 92%, p = 0.08, manual vs. 97%, p = 0.027).

The rate of severe hypoxemia (SpO2 < 80% and 85%) were reduced with FreeO2 (97%).

The rates of ventricular extrasystoles were 43, 35, 6%, p < 0.001 and atrial extrasystoles were 28, 20 and 17%, p = 0.15 for manual titration, FreeO2 (92%) and FreeO2 (97%) respectively.

The mean maximal heart rate was 96 ± 15, 86 ± 10 and 82 ± 13, p = 0.018 for manual titration, FreeO2 (92%) and FreeO2 (97%) respectively.

There was no differences in hospital length of stay between the groups and the mortality was 0% in all groups. Other data are under analysis.

**Conclusion**

Automated oxygen titration with FreeO2 with SpO2 target set at 97% reduced the rate of desaturation and severe desaturation in patients managed for acute coronary syndrome.

Automated oxygen titration with FreeO2 with SpO2 target set at 97% reduced the rate of ventricular extrasystoles.

SpO2 target set at 92% with FreeO2 may not be sufficient in this population.

With manual titration the risk of hypoxemia and severe hypoxemia was increased, however, the rate of significant hyperoxia was not increased in this study and could not be evaluated.

Additional data are required to evaluate intermediate levels of oxygenation target and potential impact in this population.

**Competing interests**

Co-founder of Oxynov company that develops the FreeO2 system.

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O60

Changes in end-tidal carbon dioxide as a surrogate for cardiac output changes outperform heart rate-, blood pressure- and femoral Doppler-derived indices during fluid challenge

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**Introduction**

During acute circulatory failure, volume expansion (VE) aims at increasing cardiac output (CO). However, CO is seldom measured to manage VE [1]. Increase in systolic, mean and pulse arterial blood pressure (ΔveSBP, ΔveMBP and ΔvePP) or decrease in heart rate (ΔveHR) are often used as surrogates for VE-induced increase in CO (ΔveCO) despite their poor performance.

VE-induced changes in end-tidal carbon dioxide (ΔveEtCO2) could be a surrogate for ΔveCO. Indeed, ΔveCO2, the amount of exhaled carbon dioxide increases with the cardiac output.
dioxide (CO₂), depends on CO₂ production by body tissues, its delivery by CO and its elimination by alveolar ventilation. If, during VE, alveolar ventilation is kept unchanged, i.e., during fully controlled ventilation, and if we hypothesize that CO₂ production only mildly changes, then ∆veCO₂, may reflect ∆veCO [2].

Other appealing surrogates were also scarcely evaluated: Doppler measurements of VE-increased in femoral artery flow (∆FemFlow) or, provided the absence of inspiratory efforts and arrhythmia, VE-induced decrease in respiratory pulse pressure variation (∆vePPV).

The objective of this study was to compare ∆veEtCO₂, ∆veFemFlow, ∆veSBP, ∆veMBP, ∆vePP and, when applicable, ∆vePPV and ∆veHR as surrogates for ∆veCO for the identification of patients who responded/did not respond to VE.

Patients and methods Adult patients were prospectively included if (1) they already had an arterial line, (2) they were receiving controlled mechanical ventilation, (3) their blood pressure (BP) was stable over 5 min, (4) the attending physician prescribed a VE and (∆) presence of at least one criterion suggesting acute circulatory failure: hypotension (invasive systolic BP < 90 mmHg and/or mean BP < 65 mmHg), oliguria (<0.5 ml kg h⁻¹) considered to be related to circulatory failure, arterial lactate >2.5 mmol L⁻¹, skin mottling, or and/or isotropic drug infusion.

CO was measured by trans-thoracic echocardiography and was the average of 2 sets of 3 (S in case of arrhythmia) consecutive measurements started from the velocity time interval of highest magnitude. EtCO₂, displayed on the ventilator, was collected at a glance, once at each study phase.

Femoral velocity time interval was measured with a 5-MHz linear echographic probe to quantify flow. Other definitions were tested for femoral flow: magnitude (peak) of the femoral pulse wave.

The ability to identify patients responding or not to VE (∆veCO > 15%) was evaluated by calculation of the areas under the receiver operating characteristic curves (AUCROCs).

Results In 109 patients included, poor thoracic insonation prevented CO measurements in 22 (20%). One patient was excluded because of change in minute ventilation >0.2 L/min during the study protocol. In the 86 remaining patients, the AUCROC for ∆veEtCO₂ was 0.82 [0.73–0.90], significantly higher than for ∆vePP, ∆veSBP, ∆veMBP and ∆veFemFlow, whatever its definition (AUCROC 0.61–0.65, all p < 0.05).

A ∆veEtCO₂ > 1 mmHg had good positive (5.0 [2.6–9.8]) and fair negative (0.29 [0.2–0.5]) likelihood ratios. Arrhythmia was of little impact on the reliability of ∆veEtCO₂: the 16 patients with arrhythmia had similar relationship between ∆veEtCO₂ and ∆veCO than regular rhythm patients (r² = 0.23 in both subgroups).

In 60 patients with no arrhythmia, ∆veEtCO₂ (AUCROC = 0.84 [0.72–0.92]) outperformed ∆veHR (AUCROC = 0.52 [0.39–0.66], p < 0.05) and tended to outperform ∆vePPV (AUCROC = 0.73 [0.60–0.84], p = 0.21).

In the 45 patients with no arrhythmia and receiving a tidal volume <8 ml/kg of ideal body weight, EtCO₂ was of significantly better performance than ∆vePPV: AUCROC = 0.86 [0.72–0.95] vs. 0.66 (0.49–0.80), p = 0.02.

Conclusion ∆veEtCO₂ outperformed ∆vePP, ∆veSBP, ∆veMBP, ∆veFemFlow and ∆veHR, and, in case of protective ventilation and/or arrhythmia, also outperformed ∆vePPV. ∆veEtCO₂ > 1 mmHg indicates that the patient is very likely to have responded to VE.

Competing interests None.

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O61 Intra-abdominal hypertension among critically ill patients requiring extracorporeal life support for refractory cardiac arrest

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Introduction Patient admitted to the ICU for refractory cardiac arrest can currently be managed using peripheral veno-arterial extracorporeal life support (ECLS) [1]. Because of impaired venous in-flow, occurrence of intra-abdominal hypertension (IAH), that is, intra-vesicular pressure (IVP) above 12 mmHg, can compromise ECLS efficiency [2]. We hypothesized that prompt diagnosis and treatment of IAH among patients under ECLS for refractory cardiac arrest may improve ECLS performance and clinical outcome. We aimed to describe critically ill patients requiring ECLS for refractory cardiac arrest and presenting with IAH.

Patients and methods This was a retrospective study of patients admitted to the ICU of the Besançon University Hospital, between January 2010 and August 2016, who develop IAH under peripheral veno-arterial ECLS for refractory cardiac arrest. We collected data about diagnosis and treatment of IAH, ECLS performance, and patient outcome. Descriptive analysis was performed. Quantitative variables were expressed as median (interquartile range) and qualitative variables as number (percentage).

Results During the study period, 202 patients (4% of the patients) were admitted to our ICU with femoral veno-arterial ECLS. Indication of ECLS was refractory cardiac arrest in 97 patients (48%), and among them, 28 (29%) developed IAH. Data were available for 27 patients who were included in the analysis. Maximum IVP was 22 [17–25.5] mmHg, which was reached 12 [5–19.5] h after ECLS onset. IVP was above 20 mmHg in 56% of cases. Curarization was initiated in 22 patients (81%). Ten patients (45%) required additional rescue laparotomy which was performed 2.5 [2–5] h after the diagnosis of abdominal compartment syndrome. ECLS parameters during maximum IAH and their improvement after efficient treatment were presented in Fig. 8. Clinically, there was a transient improvement of hemodynamic status after treatment, but hepatic and renal failures persisted. In our cohort, in-ICU mortality reached 100%. Deaths occurred after a median duration of ECLS of 32 [12–69] h, and because of refractory multi-organ failure in 82% of cases. Conclusion Among patients under ECLS for refractory cardiac arrest, IAH is frequent, occurs early, and is associated with an extremely poor prognosis. IAH treatment is associated with an improvement of ECLS outflow and hemodynamic status. IAH among patients presenting with refractory cardiac arrest might be a strong prognostic factor per se. It’s not excluded that early and efficient treatment of IAH, before occurrence of abdominal compartment syndrome, might improve the overall prognosis of critically ill patients under ECLS for cardiac arrest.

Competing interests None.
Low-dose heparin versus full systemic anticoagulation in critically ill patients undergoing extracorporeal membrane oxygenation: the HELP-ECMO pilot randomised controlled study

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Introduction Variable intensity systemic anticoagulation with unfractionated heparin is routinely used in patients undergoing extra corporeal membrane oxygenation (ECMO) to offset the increase risk of thrombosis in this population. Bleeding remains the most frequent complication and is independently associated with worse outcomes. Therefore, determining optimal anticoagulation to prevent thrombosis whilst minimising bleeding in adults on ECMO is a priority. This pilot study aims to evaluate the feasibility of allocating patients on ECMO in two different anticoagulation patterns group resulting in a difference in anti-coagulation level.

Patients and methods This is a randomised, controlled, unblinded pilot trial at two intensive care units. We enrolled critically ill patients who required ECMO (venous–venous [VV] or venous-arterial [VA]). We randomly assigned patients to therapeutic anticoagulation with heparin (target activated partial thromboplastin time [aPTT] between 50 and 70 s) or low dose heparin (12,000 units/24 h aiming for aPTT <45 s). Paired aPTT and anti-Xa assays were taken at a minimum once a day. The primary endpoints for feasibility were difference in mean heparin dose, aPTT and anti-Xa levels. All primary outcomes were log-transformed prior to analysis and reported as geometric means (95% CI) with overall differences determined using Repeat measures ANOVA.

Results Between May 2014 and March 2016, 31 patients who underwent ECMO (9 VA and 22 VV) were enrolled; 16 were randomised to low dose and 15 to therapeutic dose heparin. The groups were similar in age (mean 41 years [SD 16.8] vs 43 [SD 17.6] p = 0.75), gender (68 vs 80% male, p = 0.47), type of ECMO (31 vs 27% VA, p = 0.78) and severity sepsis-related organ failure assessment score (mean 10 [SD 3.6] vs 10 [SD 3.3], p = 1.0). The mean duration of ECMO support was 9.33 days (SD 5.97) in the low dose and 9.79 days (SD 4.77) in the therapeutic dose group (p = 0.82). For the primary outcomes, there was a significant difference in the daily mean aPTT (48.1 [95% CI 45.3–53.5] vs 56.2 [95% CI 50.7–62.3], p = 0.03), daily mean anti-Xa (0.11 [95% CI 0.07–0.18] vs 0.30 [IQR 0.19–0.46], p = 0.003) and daily mean heparin dose (11,784 units [95% CI 8693–15,972] vs 22,050 [IQR 16,262–29,899], p = 0.004) in the low dose compared to therapeutic group. There was no difference in thrombocytopenia and bleeding complications between study groups.

Conclusion In this pilot trial, administration of a low dose heparin protocol was feasible, and resulted in a significant difference in mean heparin dose administered and daily aPTT and anti-Xa levels between groups. Our findings support the feasibility of a larger study to evaluate the safety and efficacy of low-dose anticoagulation compared with therapeutic heparin with regard to thrombotic and bleeding events in patients receiving ECMO.
seem necessary to precise the best antibiotic management in these patients.

Competing interests
None.

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O67 Development and validation of a computer algorithm to detect nosocomial infections in critically ill patients

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Introduction ICU-acquired infections are common in critically ill patients and result in prolonged ICU-stay and increased healthcare costs. Moreover, the necessary antibiotic treatment may lead to a selection of multiresistant bacteria. Therefore, health authorities mandate the registration of specific nosocomial infections such as ventilator-associated pneumonia and catheter-related blood stream infections and wound infections. Unfortunately, there is no unified definition of nosocomial infection and its incidence depends on the screening intensity.

We therefore hypothesized that a computer algorithm using the prescribed antibiotic treatments may be an objective alternative for the detection of nosocomial infections.

Patients and methods As a gold standard, infections were scored by an infectious disease specialist. The Java algorithm uses the following inputs: patient identifier, admission/discharge day, antibiotic name/dose/route of administration, administration days of antibiotics. The algorithm distinguishes infections on admission and nosocomial infections.

The algorithm was trained and validated in data sets of critically ill adults (ICU) and children (PICU) admitted to the ICU of the University Hospitals Leuven between 2012 and 2015.

Results In the training sets 314/335 PICU-patients and 352/383 ICU-patients received antibiotics. Gold standard scoring yielded a nosocomial infections rate of 11% in the PICU and 16% in the ICU-population. When the criterion of at least 2 days of antibiotics beyond on admission by gold standard measurement. Here a True Positive Rate of 0.917, a False Positive Rate of 0.084, a Positive Predictive Value 0.671 and a Negative Predictive Value of 0.983 were achieved. In the validation set only adult data were used. 384/408 patients received antibiotics. 55 had a nosocomial infection and 91 an infection on admission by gold standard measurement. Here a True Positive Rate of 0.873, a False Positive Rate of 0.067, a Positive Predictive Value 0.705 and a Negative Predictive Value of 0.976 were obtained. For scoring infections on admission the algorithm had a True Positive Rate of 0.785, a False Positive Rate of 0.058, a Positive Predictive Value 0.843 and a Negative Predictive Value of 0.917.

Discussion Sensitivity (TPR) and specificity (1-FPR) are comparable and acceptable around 0.85–0.90 for an algorithm that scores nosocomial infections with limited input variables. In its current settings it could serve as elimination tool for patients who definitely do not have a nosocomial infection. Infectious disease specialists can then focus on remaining patient files for scoring nosocomial infections.

Sensitivity needs to be increased while maintaining specificity for actual use in an environment with varying antibiotic prescription practices. The algorithm should also be further validated in larger datasets for different ICUs.

Conclusion In this proof-of-concept study, a computer algorithm was shown to be an acceptable alternative to trained infectious disease specialists in scoring nosocomial infections.

Competing interests
None.

O68 Temporal trends in ICU-acquired bacteremia due to Staphylococcus aureus annual incidence in a French national ICU network

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Introduction Staphylococcus aureus (SA) is the most common isolated Gram-positive organism responsible for infection in ICU settings. The objective of this study was to assess in a French national network of ICU surveillance, the incidence of SA ICU-acquired bloodstream infection (BSI) overall and according to resistance profiles, either methicillin-susceptible (MSSA) and methicillin-resistant SA (MRSA), and the temporal trends of the incidence over 10 years, adjusted for patients’ case mix and ICUs’ characteristics.

Materials and methods Data from 2005 to 2014 were used. The incidence of BSI due to SA was assessed as the yearly ratio of ICU-acquired (>48 h.) SA BSI per 10,000 ICU patients. Only the first SA BSI occurred during one ICU stay for a patient was taken into account. Univariable autoregressive models were performed to assess the temporal trend in the evolution of the annual incidence during the study period. Multivariable autoregressive model were performed to adjust on case-mix and other centers’ characteristics with a significant annual trend selected in the univariable analysis.

Results Of 265,035 patients included from an annual median number of 158 participating ICUs (144 in 2005, 213 in 2014), 9533 (3.6%) had an ICU-acquired BSI, and 1476 (15%) of them had at least one SA BSI (56 per 10,000 ICU patients). One-third of SA BSI (n = 491, 33%) was MRSA and 907 (67%) were MSSA. There was a significant decrease of annual incidence for all SA BSI from 2005 to 2014 (48 to 10,000 ICU patients; −1.8/10,000 ICU patients per year; p = 0.02). The raw annual incidence of MRSA BSI decreased significantly from 34 to 11/10,000 ICU patients between 2005 and 2014 (−2.21/10,000 per year; p = 0.001). There was no significant change in the incidence of MSSA from 28 to 35/10,000 between 2005 and 2014 (+0.35/10,000 per year; p = 0.53) (Fig. 9). By adjusting on annual ICUs variables significant in the multivariable model (percent of medical-surgical ICU, public hospital, percentage of patients with antibiotic therapy at ICU admission, central venous catheter exposure, medical admission and annual median of age) there was no significant change in the overall incidence of SA BSI, but a significant decrease of MRSA (p = 0.02). The temporal trends were similar in all French regions.

Conclusion After adjustment on center variables, there was neither significant change in the overall incidence of S. aureus BSI nor in the incidence of MSSA BSI in ICU. However, the incidence of MRSA BSI decreased significantly. Moreover MSSA BSI did not significantly
increase. MRSA rate trends in the ICUs participating to the network (about half the French ICUs) are in line with national MRSA trends.

**Competing interests** None.

**O69** Diagnostic strategy in critically ill patients with hemophagocytic lymphohistiocytosis

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**Introduction** Hemophagocytic lymphohistiocytosis (HLH) is a life-threatening condition characterized by NK/T cells/macrophage overactivation resulting in a cytokine storm with multiorgan dysfunction. In adults, HLH is mostly secondary to cancers (mainly hematological neoplasms), infections or auto-immune diseases. HLH-etiology can be difficult to assess, especially in the critical care setting. Clinical and biological symptoms of HLH, its etiology and underlying immune deficiency being often mixed. Yet, targeting HLH-etiology is part of the ICU management in which supporting organ failure, early toposisophism and empirical anti-infectious agents are routine. In this work, we assessed diagnostic yield of tissue biopsies to identify HLH-etiology.

**Patients and methods** This single center retrospective study was conducted between 2007 and 2016. Medical records of all consecutive patients admitted with a HLH diagnosis (defined using HLH 2004 criteria) were reviewed. The performance of each diagnostic procedure was established. Rstudio software was used for analysis.

**Results** Over the study period, 142 patients (45 women, 32%; median age 47 years [37–57]) were recorded. Acute respiratory failure was the main reason for ICU admission (40 cases, 28%). Ninety-six patients (68%) had a known immune deficiency (HIV 36%, other 32%). Median SOFA at admission was 7 [5–10]. All patients met at least 5/8 HLH criteria. HLH-etiology was identified in 137 patients (96%): malignancy in 102 (74%), infection in 28 (20%), autoimmune disease in 7 (5%). Life sustaining therapy was needed in 80 patients (56%): mechanical ventilation (52%), vasopressors (44%), or renal replacement therapy (37%). Hospital mortality was 42%, with no difference between patients with or without identified HLH-etiology.

Bone marrow smears were performed in the ICU in 77 cases and were mostly useful to contribute to HLH diagnosis (83% hemophagocytosis). However, only 23 (30%) identified HLH-etiology. Among tissues that were sampled, the highest and the lowest diagnostic yields were provided by spleen resection and liver biopsy, respectively (7/8 [87.5%] splenectomies and 11/30 [37%] liver biopsies allowed to establish a definite diagnosis).

Figure 10 displays respective contributions of other tissue examination. Interestingly, the best feasibility/contribution ratio was achieved by minimally invasive lymph node biopsy. Namely, among 97 patients with clinically relevant lymph nodes, 57 (59%) could be explored under echography or CT, and diagnostic yield was 74%.

Severe adverse events included two cases of reversible hemorrhagic shocks, one following transjugular liver biopsy and one after splenectomy. No post-procedure infection was recorded.

**Discussion** Identifying HLH-etiology is a mandatory step for short and long term recovery. Moreover, when HLH patients are critically ill, identifying and targeting HLH etiology is the cornerstone of ICU management. However, in patients with multi-organ dysfunction, clinicians need guidance to understand which procedure is likely to identify HLH-etiology. In patients with HLH, pancytopenia or spleen enlarge-ment are the rule. However, other organ involvement has to be sought in each specific case.

Even if splenectomy achieves the highest diagnostic yields, minimally invasive lymph node biopsy or skin biopsy are of value based on their feasibility/diagnostic yield ratio. Only few complications were identified in this work and minimally invasive procedures seem to be safe.

**Conclusion** This study provides important guidance to establish HLH-etiology. Timing for identifying HLH-etiology is the cornerstone of ICU management. However, in patients with multi-organ dysfunction, clinicians need guidance to understand which procedure is likely to identify HLH-etiology. In patients with HLH, pancytopenia or spleen enlarge-ment are the rule. However, other organ involvement has to be sought in each specific case.

Even if splenectomy achieves the highest diagnostic yields, minimally invasive lymph node biopsy or skin biopsy are of value based on their feasibility/diagnostic yield ratio. Only few complications were identified in this work and minimally invasive procedures seem to be safe.

**O70** Granulocyte colony-stimulating factor-induced neutropenia recovery and respiratory status deterioration in critically ill patients with hematologic malignancies

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**Introduction** Granulocyte colony-stimulating factor (G-CSF) is commonly used in critically ill patients with hematologic malignancies. However, it can result in neutropenia. The aim of the study is to characterize the recovery pattern of neutropenia and the respiratory status after administration of G-CSF.

**Patients and methods** This single-center retrospective study was conducted between February and April 2016. Medical records of all patients admitted with hematologic malignancy and prescribed G-CSF were reviewed. The primary outcome was the recovery of neutrophil count ≥1500/μL. Secondary outcomes were respiratory status and mortality at discharge.

**Results** Over the study period, 30 patients (24 men, 66%) were included. Median age was 63 years (27–73), and median APACHE II score was 15 (5–28). The median duration of neutropenia was 10 days (5–30). The recovery of neutrophil count was achieved in 27 patients (90%). The median duration of recovery was 7 days (4–10). Respiratory status improved in 14 patients (46%) and worsened in 12 patients (40%). The median duration of respiratory deterioration was 3 days (1–10). One patient died during the stay due to respiratory failure.

**Discussion** Our study shows that G-CSF can induce neutropenia in critically ill patients with hematologic malignancies. The recovery pattern is rapid, and the respiratory status is rarely affected. However, further studies are needed to evaluate the long-term effects of G-CSF on neutrophil count and respiratory status.
Introduction In patients with haematological malignancies, neutropenia recovery is a situation where respiratory status may deteriorate. It has been previously demonstrated that granulocyte colony-stimulating factor (G-CSF) administration increases the risk of lung injury. ICU physicians are reluctant to use G-CSF in neutropenic patients with respiratory symptoms. However, data supporting this strategy are limited, consisting mainly of case series, retrospective data and experimental data. The aim of this study is to evaluate the impact of G-CSF administration on respiratory status in neutropenic patients with haematological malignancies admitted in ICU.

Patients and methods TRIAL-OH is a prospective, multicenter observational study that included 1011 patients with hematological malignancies who required ICU admission in 2010–2011 in 17 French and Belgian centers [1]. 288 patients who were neutropenic at ICU admission were included. The main endpoint was respiratory status deterioration at day 7, defined as any increase in respiratory SOFA score or death within the first 7 days of ICU stay. Using a propensity score (PS) based on the probability of receiving G-CSF during the first 48 h of ICU stay, we estimated the association between GCSF administration and respiratory function at day 7 on the matched sample, using a logistic regression model, adjusted on respiratory SOFA score at admission.

Results 288 neutropenic patients were included in the study. 201 (70%) did not receive G-CSF during the first 48 h of ICU stay. 87 (30%) received G-CSF at day 1 or day 2. 142 patients were selected by PS-matching. 57 (40%) were male, the median age was 58 (46–65) years. The most frequent malignancy was acute leukemia in 60 (42%) patients. The median SOFA score at admission was 6 (4–9). The respiratory SOFA score at admission was 0 for 103 patients (73%), 1–2 for 18 patients (12%) and 3–4 for 24 patients (15%). 11 (8%) patients had an invasive pulmonary aspergillosis. After propensity score matching (71 patients/group), there was no significant association between G-CSF administration and respiratory status deterioration during the following 7 days (OR 1.08; 95% CI 0.55–2.13; p = 0.83), even though neutropenic patients who received G-CSF had better neutrophil recovery at day 7. 70 (49%) patients died or showed an increase of the respiratory SOFA score. Among the 37 (53%) respondents, 33 (47%) reached the sensitivity analysis in patients admitted for acute respiratory failure, we estimated the association between GCSF administration and respiratory status deterioration during the following 7 days (OR 1.08; 95% CI 0.55–2.13; p = 0.83). No adverse reaction was noted following Rituximab infusion during the ICU period. Follow up was 1126 days (271–2001) and 3 additional patients died during follow-up. Eleven (14%) patients had TMA recurrence within 585 days (125–858). There was a trend toward a longer time to recurrence in Rituximab-treated patients (1475 days [427–2530] vs 136 days [80–749], p = 0.09). In addition, 22 TTP patients subsequently received Rituximab during follow-up either for TTP clinical recurrence or as maintenance therapy for TADMA. No adverse reaction was noted following Rituximab infusion during the ICU period. Follow up was 1126 days (271–2001) and 3 additional patients died during follow-up. Eleven (14%) patients had TMA recurrence within 585 days (125–858). There was a trend toward a longer time to recurrence in Rituximab-treated patients (1475 days [427–2530] vs 136 days [80–749], p = 0.09). In addition, 22 TTP patients subsequently received Rituximab during follow-up either for TTP clinical recurrence or as maintenance therapy for persistent or recurrent ADAMTS13 undetectable activity. Long term complications potentially attributable to Rituximab included transient decrease in IgM levels (3 patients), serum disease in 1 patient, mild cognitive dysfunction in 1 patient and asymptomatic hepatitis B virus reactivation in 1 patient.

Discussion Rituximab, when used in patients with severe/refractory TTP, seems safe at short and long term. This apparently low complication rate may be due to the relatively low Rituximab dose regimen as compared to regimen applied to patients with lymphoma. High dose steroids might also have been protective against acute cytokine reactions. Last, the retrospective nature of this study might also have undermined transient or mild Rituximab-related toxicity.
Conclusion In critically ill patients with severe/refractory TTP, safety profile of Rituximab is good. Studies to assess its efficacy as early adjunctive therapy in non-refractory TTP patients are warranted.

Competing interests None.

O72 Long term health related quality of life in critically ill patients with hematological malignancies

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Introduction Outcome of critically ill patients with hematological malignancies has substantially improved over the last decade. However, data regarding the long term Quality of Life (QOL) of these patients are scarce. The purpose of this study is to assess post ICU discharge related QOL, and to identify risk factors for long term QOL impairment.

Patients and methods TRIAL-OH is a prospective, multicenter observational study that included 1011 patients with hematological malignancies who required ICU admission in 2010-2011 in 17 French and Belgian centers. Ninety days and 1 year after ICU discharge, HRQOL was was determined by applying the interview form of the short-form 36 questionnaire (SF-36), which measures HRQOL in 8 separate dimensions (0 = worst health state, 100 = best health state). Psychological distress symptoms were evaluated using the Hospital Anxiety Depression Scale (HADS) and the Impact of Event Scale (IES). All data were collected prospectively.

Results Two hundred and seventy-nine patients were evaluated at 3 months and 117 patients were evaluated at 1 year after ICU discharge. At 3 months global median Physical (PS) and Mental scores (MS) were respectively 51.6 (48.0–55.3) and 49.7 (46.1–53.2), and at 1 year median PS and MS were respectively 63.8 (59.9–67.6) (p = 0.92) and 56.3 (52.5–60.0) (p = 0.05).

Physical functioning, general health, social functioning, mental health did not significantly change between 3 months and 1 year. Role Limitation due to physical problems (0.0 (0–6.2) vs 25.0 (17.7–32.3), p = 0.0008) and Vitality Score (40.0 (36.1–43.9) vs 46.7 (43.3–50.0) p = 0.018) significantly improved at 1 year. Role limitation due to emotional problems (72.4 (93.4–106.6) vs 66.7 (59.2–74.1), p = 0.001) and bodily pain (70.0 (64.2–75.8) vs 67.5 (63.0–72.0) were significantly worse at 1 year. At 3 months 69 (17.5%) patients had an IES score at 20 and 139 (49.5%) patients had a HADS at 8.

Discussion This multicentric study is the first to assess long term QOL in a large cohort of critically ill patients with hematological malignancies. Significant improvement in terms of short term mortality in this population do not fully reflect the impact of ICU stay.

Prompt recognition of risk factors for impaired QOL and adapted therapy for anxiodepressive symptoms may improve their outcome.

Conclusion Critically ill patients with hematological malignancies have impaired QOL at 3 months and recovery is incomplete 1 year after ICU discharge. Long term QOL should be taken into account in the management of these patients.

Competing interests None.

O73 Should we admit patients with hematological malignancies earlier to the ICU?

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Introduction Over the last two decades, advances in hematology and critical care management have translated into improved outcomes for critically ill hematological patients. Concerns have been raised about delayed admission to the ICU, however, the strength of evidence remains weak.

Patients and methods In a substudy from Triahol led by the GRRR-OH and published in 2013, we sought to assess how time from hospital to ICU admission is associated with hospital mortality. Data were collected from 17 ICUs from France and Belgium between January 2010 and May 2011. We used non parametric Wilcoxon tests and Fisher exact tests for baseline univariate comparisons across groups, p-value equal or below 0.05 were considered statistical significant. Regression logistic models were used to summarize predictive information for ICU mortality.

Results Based on non-parametric estimation, the cutoff value of 3 days for time between hospital and ICU admission was best associated with mortality (OR 1.66 (1.28–2.15), P = 0.0001). Among the 1007 included patients, 592 were admitted before day-3 and 415 after day-3. Non-Hodgkin’s lymphoma, acute myeloid leukemia, and myeloma represented about 70% of admissions in the two groups. Patients admitted earlier were older (62 (50–71) vs. 59 (48–67), P = 0.01), more frequently women (35 vs. 45%, p = 0.003), had higher Charlson scores (4 (3–6) vs. 3 (2–5), P = 0.007 mostly from diabetes, cardiovascular diseases and chronic kidney diseases), but less altered performance status 1 (0–2) vs. 2 (1–3), P < 0.001. Early admitted patients, had less frequently admitted to the ICU from the ED/SAMU (43 vs. 2%, p < 0.0001) and less frequently enrolled in a hematology trial (23 vs. 33%, p = 0.0004). Patients admitted before day-3, more frequently carried a newly diagnosed malignancy (28.69 vs. 15.74%, p < 0.0001) and were less frequently acute myeloid leukemia patients (23 vs. 33%, p = 0.0009), recipients of hematopoietic stem cell transplants (both for autologous (18.16 vs. 11.53%, p = 0.004) and allogeneic (20.29 vs. 10.34%, p < 0.001)) or patients not in remission (36 vs. 46%, p < 0.0001). Delayed ICU admission occurred more frequently in...
patients with acute respiratory failure (70.3 vs. 57.2%, p < 0.0001) or coagulation disorders (23.7 vs. 16%, p = 0.003). Patients admitted after day-3, developed more frequently multiple organ failure (59.5 vs. 51%, p = 0.009).

By multivariable analysis with mortality as the outcome variable, admission after day-3 tended to be associated with mortality but did not reach significance when adjusted on patient's characteristics and processes of care. An ongoing analysis to adjust on severity at ICU admission will be communicated in January.

Conclusion Patients admitted to the ICU more than 3 days after hospital admission have an increased mortality. As one-third of these patients were admitted after an initial refusal, these results identify the typology of patients for whom delayed ICU admission impacts on outcomes. Interventional studies are warranted.

Competing interests None.

O74 The effects of passive leg raising can be detected by the plethysmographic oxygen saturation signal

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Introduction Volume expansion is aimed at increasing cardiac output. Nevertheless, a cardiac output monitoring technique is not always available. The perfusion index (PI) (Masimo Corp., Irvine, CA, USA), is automatically calculated from the plethysmographic waveform of oxygen saturation as the ratio of the pulsatile fraction, caused by blood flow, and the non-pulsatile fraction. We hypothesized that PI could be proportional to stroke volume and that it could detect changes in cardiac output during passive leg raising (PLR) and volume expansion (VE).

Patients and methods We included patients for which a PLR test was planned. We measured the changes in cardiac index (CI, PiCCO2 device, Pulsion Medical Systems, Munich, Germany) and PI before and during the PLR tests. If a VE (500 mL of saline infusion over 10 min) was performed, we also measured its effects on CI and PI.

Results Fifty-five PLR tests were performed in 30 patients. One case was excluded because of a poor oxygen saturation signal. Norepinephrine was administered in all cases at a mean dose of 1.6 ± 1.0 μg/kg. The PLR test was positive (increase in CI ≥ 10%) in 26 “preload responsive” cases and negative in 29 “preload unresponsive” cases. During PLR test, in preload responsive cases, CI significantly increased by 30 ± 15% and PI significantly increased by 79 ± 45%. During PLR test, in preload unresponsive cases, neither CI nor PI changed significantly. PI was able to detect a positive PLR test with good accuracy (area under the receiver operating characteristic curve: 0.99 [95% confidence interval 0.90–1.00, p < 0.001]). If PI increased >23%, a positive response to PLR could be diagnosed with a sensitivity of 100% (84–100%) and a specificity of 90% (74–98%). Volume expansion was administered in 15 cases with a positive PLR test. Taking into account both PLR and volume expansions, the changes in CI and PI (27 ± 11% and 73 ± 47%, respectively) were correlated (r = 0.73, p < 0.001). The PI value at baseline was <2 in 15 patients, but the ability of PI to track changes in CI was not poorer in these patients than in the other ones.

Conclusion The results of this preliminary study are that the perfusion index using pulse oximetry seems to accurately reflect changes in CI during PLR test and volume expansion. This could be a reliable way to assess preload responsiveness in a totally non-invasive way.

Competing interests JLT and XM are members of the Medical Advisory Board of Pulsion Medical Systems. JLT gave lectures for Masimo. The authors have no other conflicts of interest to declare.

O75 Exploration of tissue perfusion parameters around tracheal intubation procedure and mechanical ventilation initiation

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Introduction Mechanical ventilation is a common organ support therapy in critically ill patients. Mechanical ventilation initiation could have several hemodynamic consequences, ranging from tachycardia to life-threatening collapse. The objectives of this study were (i) to describe the microcirculatory and macrocirculatory changes following emergency tracheal intubation (TI) and (ii) to identify predictors of hemodynamic instability.

Patients and methods Prospective observational monocenter study conducted in a 18-bed medical ICU. Consecutive patients requiring tracheal intubation were eligible for the study. Data collection included general demographic parameters and comorbidities, indication for tracheal intubation and presence of sepsis. Global hemodynamic parameters (blood pressure, heart rate, cardiac index) and tissue perfusion parameters (mottling score, capillary refill time [CRT], toe-to-rectal gradient temperature) were recorded before, 5 min and 2 h after TI. Cardiac function parameters (left ventricular ejection fraction, right ventricular to left ventricular diameter ratio, presence of paradoxical septum), arterial blood gas and arterial lactate level were measured before and 2 h after TI. Hemodynamic instability requiring intervention (HII) was defined as any hemodynamic status degradation requiring intravenous fluid resuscitation (≥500 ml) or the introduction of vasopressors or an increase in vasopressor dose ≥20%. Difficulty of tracheal intubation was assessed using the Intubation Difficulty Scale.

Results Seventy-seven patients were included (male gender 61%) with a median age of 69 [interquartile range 56–81] years. Median SOFA score and SAPS2 were 7 [4–10] and 51 [41–64], respectively. Indication for tracheal intubation was hypoxemia (44%), hypercapnia (14%), and coma (32%). Most of the patients had sepsis (64%), including septic shock patients (22%).

Mechanical ventilation had no significant impact on cardiac index, heart rate and tissue hypoperfusion, whereas median mean arterial pressure decreased from 82 [71–93] mmHg before intubation to 73 [66–84] mmHg 2 h after TI (P = 0.005, Wilcoxon signed rank test). HII occurred in 38 patients (48%). Univariable comparison (Fisher’s exact test for discrete variables and Mann–Whitney U test for continuous variables) of these patients with those who did not experience HII indicated that male gender (P = 0.035), intubation for hypoxemia (P = 0.006), administration of norepinephrine before TI (P = 0.002), sepsis (P < 0.0001), higher SOFA score (P = 0.0005), higher SAPS2 (P = 0.006), mottling score ≥3 before TI (P = 0.046), higher baseline serum lactate level (P = 0.038) and higher baseline knee CRT (P = 0.029) were predictors for HII. In contrast, cardiac index, mean arterial pressure, baseline temperature gradient, index CRT before TI and Intubation Difficulty Scale were not associated with HII.

Sepsis patients had a relative risk (RR) of HII of 2.8 [95% confidence interval 1.8–4.4]. Among sepsis patients, a mottling score ≥3 was associated with a RR of HII of 9.8 [1.3–73.4], serum lactate level with a RR of 2.6 [1.2–5.6], and intubation for hypoxemia with a RR of 2.1 [1.2–3.5], as compared with the entire study population.

Conclusion In a non-selected ICU population, half of the patients required fluid resuscitation or vasoactive drugs in the 2 h following TI. Sepsis was the strongest predictor of hemodynamic instability. Tissue hypoperfusion parameters, especially mottling score, identified a subgroup of septic patients with a high risk of hemodynamic instability. In contrast, global hemodynamic parameters such as cardiac index before TI did not predict bad hemodynamic tolerance of the procedure.

Competing interests None.
**O76**

The GRACE risk score in critically ill patients with sepsis and a suspected myocardial infarction

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3Annals of Intensive Care 2017, 7(Suppl 1):O76

**Introduction**

Elevations of cardiac troponin values, suggesting a myocardial necrosis, are common in critically ill patients with sepsis and are associated with a poor prognosis. In this context, myocardial ischaemia is one of the main underlying mechanisms leading to necrosis of myocardial cells defining the myocardial infarction (MI). However identifying which septic patients with suspected myocardial infarction are at risk of mortality and major cardiac events is a clinical challenge [1]. Clinical evaluation may lack sufficient precision, leading to inappropriate medication and discharge. It is uncertain whether risk scores derived from cardiological populations apply in this context. We aimed to assess whether the «Global Registry of Acute Coronary Events» (GRACE) score may predict mortality in the intensive care unit (ICU) for septic patients with a suspected MI.

**Patients and methods**

We conducted a prospective monocenter observational study from June 2012 to August 2016 in the medico-surgical ICU of Tenon Hospital, Paris, France. All patients with a suspected MI [significant cardiac troponin elevation with at least one of the following: symptoms of ischaemia, new significant ST-segment or T wave changes at electrocardiogram, acute left ventricular systolic dysfunction (1)] on ICU admission or during their ICU stay, were screened. Patients admitted for sepsis were included. The primary endpoint was to assess the performance of the GRACE score to predict ICU mortality. The secondary endpoints were to describe the respective occurrence of major cardiovascular events (stroke, cardiac arrest and reinfarction), major bleedings and cardiogenic shock during ICU stay.

**Results**

During the study period, 238 out of 3774 patients (6.3%) had a suspected MI. Among them, 122 (51%) were admitted for sepsis (75% of suspected myocardial infarction occurred at admission). Among them, 122 (51%) were admitted for sepsis (75% of suspected MI). The ICU length of stay was 8 days (4–15). The ICU mortality rate was 0.30 (95% CI 0.22–0.39). At the time of suspected MI, the GRACE score was higher in the non-survivor group (201 [160–226]), as compared with the survivor group (176 [149–211]; p = 0.053). A cut-off value of 200 was associated with ICU mortality (OR 2.53, 95% CI 1.14–5.61; p = 0.022). The symptoms of ischaemia (13 vs 8%, p = 0.441), the level of high sensitivity cardiac troponin peak (1093 pg/mL [381–4054] vs 1248 pg/mL [554–2973], p = 0.66), and the left ventricular ejection fraction at echocardiography (29% [20–41] vs 35% [20–50], p = 0.21) were similar between non-survivors and survivors. The electrocardiogram ST-segment or T wave changes were less frequent in the non-survivors than in the survivors (50 vs 69%, p = 0.041). Major cardiovascular complications, major bleedings, and cardiogenic shock occurred more frequently in the non-survivors than in the survivors (19 vs 4%, p = 0.012; 16 vs 7%, p = 0.086; 76 vs 36%, p < 0.001, respectively).

**Discussion**

Critically ill septic patients with suspected MI had a high risk of cardiovascular events and bleeding. In this population, the GRACE scoring system was more accurate for predicting ICU death than symptoms of ischemia, troponin elevation, and left ventricular dysfunction. The systematic application of a validated risk score to septic ICU patients with a suspected MI could enable an early appropriate management, and provide a more reliable basis for adequate discharge and follow-up. These findings require validation in larger prospective studies.

**Conclusion**

The GRACE risk score could be a promising score to help for predicting death in septic ICU patients with a suspected myocardial infarction.

**Competing interests**

None.

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**O77**

Systemic capillary leak syndrome severe attacks admitted in intensive care unit

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Annals of Intensive Care 2017, 7(Suppl 1):O77

**Introduction**

Systemic Capillary Leak Syndrome (SCLS) is a rare disease characterized by recurrent life-threatening attacks of capillary hyper permeability in the presence of a monoclonal gammapathy (MG). During acute episodes, the leak of fluid and proteins from the capillary compartment to the interstitium results in clinical signs of both acute hypovolemia and interstitial edema. Biological profile is pathognomonic with marked hemoconcentration and paradoxal hypoproteinemia. There are few data available about natural history and prognosis of severe SCLS attacks. The objectives of this study were to precisely describe natural history, outcome and mortality associated factors in severe SCLS attack.

**Patients and methods**

Multicenter retrospective analysis of data from the European Clarkson registry (EuréClark). Criteria to retain SCLS’s diagnostic were: presence of a MG; ≥1 typical attack with clinical manifestations of hypovolemia and capillary leak; hemoconcentration with paradoxal hypo proteinemia; exclusion of secondary capillary leak syndrome causes. Patients with severe attacks admitted in ICU were identified in EuréClark registry. Physician were contacted and performed a structured clinical interview.
offered to include the attack using a pre-established case report form. Categorical variables are expressed as n (%) and continuous variable: mean ± SD or median [IQR].

**Results** Between May 1992 and February 2016, 59 attacks in 37 patients were included. Sex ratio was 1.05 with an age of 51 ± 11.4 years. In ten patients, more than one attack was included. Thirty-four (92%) patients had an IgG MG with Kappa light chain in 20 (59%) patients. A trigger of the attack could be found in 34 (58%) patients with flu-like syndrome being the most frequent (89%). SAPS II score at admission was 54 (38–67) and SOFA score 6 (3–9). Admission heart rate was 128 ± 21 bpm, admission arterial systolic, mean and diastolic blood pressure were respectively 75 [55–94], 60 [44–70] and 45 [36–60] mmHg. Frequent clinical manifestations at admission were: unaltered consciousness despite profound arterial hypotension (83%), asthenia (78%), faintness (64%), nausea and vomiting (53%), edema (51%), dyspnea (46%), marbles (42%), myalgia (39%) and abdominal pain (36%). Admission hemoglobin, proteinemia, serum creatinine and arterial lactate were respectively 20.2 [17.9–22] g/dL, 50 [36.5–58.5] g/L, 176 [121–244.5] μmol/L and 4.6 [3.3–6.5] μmol/L. Five patients underwent phlebotomy for mistaken hyper viscosity syndrome and 57 (97%) received fluid therapy with a cumulated volume of 4.5 [2.8–10.6] L over 1 [1–2.5] days. Norepinephrine was administered in 28 (47.5%) patients, epinephrine in 10 (17%) and corticosteroids in 19 (32%). Twenty-two (37%) patients required mechanical ventilation and 57 (97%) underwent renal replacement therapy. Fifteen patients (25%) were treated with intravenous immunoglobulins (IgIV) during the attack. Compartment syndrome occurred in 12 (32%) patients and 11 (30%) died in ICU. In unvariable analysis (over 37 unique patients) main factors associated with mortality were SAPS II (p = 0.006) and SOFA score (0.005), neurological dysfunction (neuro SOFA score > 3, p = 0.003), high ICU maximum weight (p = 0.008), high cumulated volume of fluid therapy (p = 0.017), mechanical ventilation (p < 0.001), and renal replacement therapy (p = 0.002) but not treatment with IgIV (p = 0.12). Multivariable analysis retained a SOFA > 10 (OR 10.4 [1.1–91], p = 0.04) and a cumulated volume of fluid therapy > 8 L (OR 16.4 (1.2–230), p = 0.03) as independent factors of mortality. Treatment with IgIV (OR 11.5 [0.85–155], p = 0.06) was not independently associated with mortality.

**Conclusion** Our study presents the first large cohort of SCLC attacks included to ICU. Compartment syndrome was a particularly frequent complication and mortality over 59 attacks was 18.6%. High cumulated volume of fluid therapy seems to be associated with poorer outcome. There was a trend towards mortality in patients treated with IgIV during the attacks and such treatment should be used with caution.

**Competing interests** None.

**SHORT PRESENTATIONS**

**P1 PHARMECMO: a pilot pharmacokinetic study of antibiotics in patients assisted by extracorporeal life support**

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The PHARMECMO study was a pilot, prospective, pharmacokinetic study, conducted in ICU cardiac surgery. All patients treated with ECLS, with known or suspected sepsis and receiving antibiotic therapy, were eligible for inclusion. The objective of this pilot study was to observe the pharmacokinetic characteristics of commonly used antibiotics in intensive care unit (ICU) for patients treated with ECLS.

**Patients and methods** The PHARMECMO study was a pilot, prospective, pharmacokinetic study, conducted in ICU cardiac surgery. All patients treated with ECLS, with known or suspected sepsis and receiving antibiotic therapy, were eligible for inclusion. The concentration of each antibiotic was measured by a combination of liquid chromatography and mass spectrometry from blood samples. For intermittent administration of antibiotic, two successive measures were performed, under steady state conditions, both at 50% (CT50) and 100% (Cmin) of the dosing interval.

**Results** Forty-five eligible patients were enrolled for 68 inclusions allowing 114 analysed samples during 1 year. The median age was 63 years (interquartile range [IQR] 58–67), 84.4% of inclusions allow 114 analysed samples during 1 year. The median age was 63 years (interquartile range [IQR] 58–67), 84.4% of inclusions
were male, with a median weight of 74.5 kg (IQR 70–86.75). Among the 68 inclusions, 39.7% received continuous veno-venous hemofiltration. ECLS therapy was veno-venous (n = 2), peripheral venoarterial (n = 51) or central venoarterial (n = 15). The most frequent causes of infection were pneumonia (n = 36), infection of femoral triangle (n = 5) and catheter-associated infection (n = 5). Of the pathogens identified, *Pseudomonas aeruginosa* was the most frequent (n = 20).

The main studied antibiotics were piperacillin-tazobactam (n = 17), cefotaxime (n = 12), imipenem (n = 10) and amikacin (n = 6). For the association piperacillin-tazobactam, the median CT50 was 87.00 mg/L (IQR 57.93–158.71) and the median Cmin 61.24 mg/L (IQR 44.92–90.51) for a dose of 4 g four times a day, with a MIC target of 16 mg/L. For cefotaxime, median concentrations were respectively, CT50 and Cmin of 64.69 mg/L (IQR 20.17–97.52) and 28.61 mg/L (IQR 7.28–42.12) for a MIC target of 1 mg/L and a median dose of 7 g per day. Regarding imipenem, at a dose of 1 g three times a day, the median concentrations were respectively, CT50 and Cmin, of 7.30 mg/L (IQR 4.07–14.59) and 3.28 mg/L (IQR 1.84–5.43) for a MIC target of 4 mg/L. Only one patient had a CT50 greater than 4 MIC, and 60% of measured Cmin were under the MIC. Finally, for amikacin, the median Cmax was 51.14 mg/L (IQR 32.36–78.26) at a medium dose of 24.3 mg/kg, for a target between 60 and 80 mg/L.

Conclusion These preliminary data suggest that therapeutic drug monitoring could optimize the achievement of pharmacokinetic objectives associated with an effective antibiotic therapy. These data also suggest that, in most patients, the recommended doses of imipenem at 1 g three times a day and aminoglycoside at 20 to 25 mg/kg, do not respect the PK objectives reported in the literature.

Competing interests None.

### P2

**Variation in screen and isolate policy for multidrug-resistant bacteria (MDRB): a national survey in French adult ICUs**

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**Annals of Intensive Care 2017, 7(Suppl 1):P2**

**Introduction** The control of health-care associated infections and multidrug-resistant bacteria (MDRB) is a public health priority. The MDRB prevalence rate has dramatically increased, mainly extended-spectrum beta-lactamase producing Enterobacteriaceae (ESBLE). The aim of this study is to describe and to analyse the different screen and isolate policies regarding MDRB in French adult ICUs.

**Patients and methods** This is an observational, descriptive, multicenter online survey performed in French adult ICUs. The questionnaire included 63 questions divided into 4 parts: characteristics of the unit, MDRB screening policy, policy regarding contact precautions and methods. Imported and acquired MDRB rates >10% were noted in 44% and 27% of cases, respectively. Almost half of the units (48%) had already experienced a MDRB epidemic situation in the 3 preceding years.

**Results** From April 2015 to June 2016, 73 of 301 French adult ICUs responded to the survey (24% response rate). A screening upon admission was performed in 96% of cases, mostly in a systematic way for at least one MDRB (78%). Screening was commonly used for admission was performed in 96% of cases, mostly in a systematic way (71%). Participants varied on gown and gloves wearing from standard precautions to consolidated additional contact precautions. About 23% of units used one or several specific decontamination methods. Imported and acquired MDRB rates >10% were noted in 44% and 27% of cases, respectively. Almost half of the units (48%) had already experienced a MDRB epidemic situation in the 3 preceding years.

**Discussion** French adult ICUs vary significantly in their MDRB screening and isolate approach, and about 10 combinations were encountered in the survey. The different approaches practiced were not always in agreement with the 2009 national guidelines for the prevention of MDRB transmission. Very few ICUs proceeded without screening and isolation at admission.

**Conclusion** Substantial variations exist in French ICUs practices regarding MDRB screening and isolation. The growing impact of imported and acquired MDRB rates and the frequency of epidemic situations in the ICUs may induce to reconsider and clarify the recommendations for prevention of “cross-transmission” of the French Society of Hospital Hygiene.

### P3

**Impact of routine decontamination on *Pseudomonas aeruginosa* acquired infections and antimicrobial susceptibility in an ICU**

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**Annals of Intensive Care 2017, 7(Suppl 1):P3**

**Introduction** Selective decontamination with various regimes using topical antimicrobials has been reported to reduce acquired infections (AIs) and mortality in the ICU. The efficacy of decontamination on the prevention of *P. aeruginosa* AIs is controversial [1] and potential impact on the emergence of antimicrobial-resistant strains is a concern. We assessed the incidence of *P. aeruginosa* AIs (PAAs) with the routine use of a multiple decontamination regimen with oropharyngeal and digestive tobramycin/colistin/amphotericin B and nasal mupirocin/chlorhexidine body wash over 5 years with a special attention to antimicrobial resistance.

**Patients and methods** This was an observational single center study of all patients admitted to an ICU 2008–2012 (study population). Decontamination was given for the period of intubation and standard care otherwise. PAAs were prospectively recorded. Rates of PAAs (proportion of patients) were first compared between the study period and a 4-year pre-intervention period (2003–2006). During study, trends were analyzed by year using a logistic regression model. Categories were compared by Chi square test or Fisher’s exact test when appropriate. Continuous variables were expressed by median (25th–75th percentile) and compared using non-parametric test.

**Results** Of the 5250 patients admitted to the ICU during the 5-year study period, 69 (1.3%) acquired 77 episodes of PA infection (vs 112 of 3603 patients [3.1%] during the pre-intervention period, p < 0.001). The incidence of PAAs declined over time (OR 0.81 [0.68–0.96], p = 0.02). The proportion of patients who had clinical samples positive for PA at admission was 1.3% (70/5250) and did not vary with time (OR 1.03 [0.87–1.22], p = 0.75). Baseline characteristics of PAAs patients were: SAPS II 55 (44–68), SOFA score 9 (7–12), prior antimicrobials exposure (number of molecules): 4 (2–5). Alts sites (n = 80) were respiratory (n = 52, ventilator-associated in 50), bloodstream (n = 15), genitourinary (n = 5), abdominal (n = 4) and other (n = 4). The delay of onset after admission was 13 days (7–22). Prior decontamination duration was 10 days (5–18). The susceptibility rate to 10 antimicrobials of the 80 isolates tested was as follows: ticarcillin (66.3%); piperacillin/tazobactam (82.5%); ceftazidime (93.8%); imipenem (52.5%); gentamicin (57.3%); tobramycin (97.5%); amikacin (96.3%); ciprofloxacin (67.5%); fosfomycin (61.3%); colistin (98.8%). Susceptibility to all antimicrobials remained unchanged with time except for fosfomycin (susceptibility rate increased with time, OR 1.59 [1.05–2.39], p = 0.03). Antimicrobial therapy of 77 episodes consisted of a β-lactam agent (89.6%), ciprofloxacin (36.3%), aminoglycoside (31.1%), 6 episodes were not treated. The mortality rate in ICU was similar in the patients with PAAs (30.4%) and in those (n = 166) who acquired AIs not due to PA (31.3%).
adjusted odds ratio = 0.84 [0.44–1.57], p = 0.58). The ICU mortality rate differed according to the site of the first episode: bloodstream 5/12 (41.6%); lung 16/46 (34.8%); other 0/11 (p = 0.03). In the patients with bloodstream or respiratory PAAIs, in multivariate analysis, SAPS II (per one unit increase OR 1.05 [1.00–1.10]) and non-susceptibility to piperacillin/tazobactam (OR 5.46 [1.04–27.77]) were independent risk factors for death in ICU (both p < 0.05). Aminoglycoside combination therapy was not associated with a higher cure rate (41%, vs no aminoglycoside 63%; p = 0.12).

Discussion Although all-cause AIs are declining in our ICU and the incidence rate has become low [2], PA remains the most common agent of AIs. 65% of PAAIs were ventilator-associated pneumonia. PAAIs were not associated with a higher ICU mortality rate than AIs due to other organisms.

Conclusion With the routine use of a decontamination regimen, AIs involving PA were controlled as well with no increase in antimicrobial resistance.

Competing interests None.

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P4 Penicillin G susceptibility among Staphylococcus aureus is not so infrequent: a retrospective study
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Introduction The incidence of resistance of Staphylococcus aureus to methicillin has decreased in France during the past decade leading to an increase use of methicillin and antistaphylococcal cephalosporins. It is usually stated that about 5% of methicillin-susceptible Staphylococcus aureus (MSSA) isolates are sensitive to penicillin G and penicillin G is infrequently used to treat Staphylococcus aureus related infection. The aim of this retrospective observational study was to assess the rate of susceptibility to penicillin G among clinical Staphylococcus aureus strains isolated from ICU patients.

Patients and methods For a 10 years period, methicillin and penicillin susceptibilities of all Staphylococcus aureus strains isolated from clinical samples were analyzed. Repeat episodes and screening samples were excluded. Demographic data (age, sex), SAPSII score and in-ICU survival were recorded and analyzed according to penicillin G susceptibility.

Results From 01/2006 to 12/2015, 584 Staphylococcus aureus strains were isolated, 136 of whom being obtained from blood cultures, 327 from respiratory samples, 25 from urines 8 from arthrosis. One hundred and three (17.6%) were susceptible to penicillin. Among all strains of Staphylococcus aureus isolated, the annual incidence of penicillin-susceptible strains varied from 5 to 24%. Over this period, global methicillin-resistant strains incidence was 15% with annual average rate ranging from 4 to 31%. No difference was observed between age, sex of patients, SAPSII score or prognosis according to penicillin-G susceptibility (Table 8).

Conclusion This retrospective study suggests that penicillin susceptibility among clinical isolates of Staphylococcus aureus is not negligible and more frequent than the classical 5%. Microbiological laboratories should continue to test penicillin on Staphylococcus aureus strains. In this era of oxacillin shortage, penicillin G could be an appealing alternative in about 20% of cases.

Competing interests None.

P5 Impact of changing third-generation cephalosporin policy use on multiple-drug resistant bacteria nosocomial infections rates
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Annals of Intensive Care 2017, 7(Suppl 1):P5

Introduction Multiple-drug resistant bacteria carriage in general population and nosocomial infections are increasing in France and Europa even efficiency of hand hygiene and antibiotic stewardship to control major hospital pathogens. Otherwise, antibiotics use have different consequences on bowel bacteria selection pressure. High biliary excretion of ceftriaxone and its long acting properties are responsible for a higher selection pressure in comparison with cefotaxime, an antibiotic drug with the same microbial spectre, lower biliary excretion, shorter action and a negligible effect on the bowel microflora.

Patients and methods Consequently, a new antibiotic policy was implemented in a 350-bed general hospital in Lyon (France) with the promotion of cefotaxim use instead of ceftriaxone from 1st January 2014 after an educational campaign in December 2013. Hospital-acquired (HA) bloodstream meticillin-resistant Staphylococcus aureus (MRSA), third-generation cephalosporin-resistant Enterobacteriaceae (3GCRE), Candida infections and Clostridium difficile colitis cases per month/1000 patient-beds were identified and reviewed throughout our 12 beds-intensive care unit (ICU).

Monthly consumption of several antibiotics was monitored in defined daily doses (DDD) per 1000 patient-occupied bed-days (1000 pt-bds) 24 months before (2012–2013) until 24 months after new antibiotic policy introduction (2014–2015).

Results Physicians have quickly followed new antibiotic policy and average monthly consumption of ceftriaxone reduced by 82% (from 119.6 to 26.6 DDDS/1000 pt-bds, p < 0.001) and cefotaxime increased in 40% (from 27.7 to 68.8 DDDS/1000 pt-bds, p < 0.001) between the first and final 24 months of the study in ICU. Over the same periods, HA bloodstream infections rates were stable between the period before and after intervention for 3GCRE (1.71 vs 1.10 cases/1000 pt-bds, p = 0.33), MRSA (0.24 vs 0.08 cases/1000 pt-bds, p = 0.41), Candida (2.09 vs 2.76 cases/1000 pt-bds, p = 0.45). And HA C difficile infections were similar between the two periods (0.60 vs 0.67 cases/1000 pt-bds for the second period, p = 0.82) in ICU.

Table 8 Age, sex ratio, SAPSII score and in-ICU survival according to PeniG susceptibility

| Age | Male | Female | SAPSII | Survival |
|-----|------|--------|--------|----------|
| 64 ± 17 | 58% | 42% | 48 ± 21 | 70% |
| 64 ± 18 | 58% | 42% | 49 ± 19 | 70% |
| 66 ± 17 | 60% | 40% | 48 ± 2 | 69% |

| N | N = 584 | N = 103 | N = 481 |
|---|---------|---------|---------|
| Age | 64 | 64 | 66 |
| Male | 58% | 58% | 60% |
| SAPSII | 48 | 49 | 48 |
| Survival | 70% | 70% | 69% |
Hand-hygiene compliance with alcoholic formulation (7.9 vs 10.31 ml per patient-days, $p = 0.12$) and consumption of all antibiotics (3936 vs 4520 DDDs/1000 pt-bds, $p = 0.12$) haven’t significantly changed before and after new antibiotic policy in our ICU and hospital.

**Discussion** While multiple-drug resistant bacteria nosocomial infections dramatically increase in France and Europe, trends in bloodstream 3GCRE, MRSA, *Candida* infections and *Clostridium difficile* colitis rates were stable in our ICU 2 years after introduction of a new antibiotic policy for third-generation cephalosporins. And our results seem different to some previous studies because only bloodstream infections were considered in order to affirm multiple-drug resistant bacteria infections. But this method induced lower HA infections rates. Our results have to be analyzed after adjustment with multiple-drug resistant bacteria carriage at admission in ICU between the two periods. About 20% of patients carriers of multiple-drug resistant bacteria will be infected by the same bacteria during hospitalization indeed. A Poisson regression model (adjusted for overdispersion) will be used with log occupied bed-days as an offset. Linear trend terms were used to assess temporal changes. The main hypothesis was to see whether the trend in HA infections rates following antibiotic policy change was the same as the trend before the intervention. Poisson regression will be again used to assess which of the antibiotics was the better predictor of HA bloodstream 3GCRE, MRSA, *Candida* infections and *Clostridium difficile* colitis rates.

**Conclusion** Hospital new antibiotic policy promoting cefotaxime use instead of ceftriaxone didn’t seem to impact hospital-acquired bloodstream meticillin-resistant *Staphylococcus aureus*, third-generation cephalosporin-resistant Enterobacteriaceae and *Candida* infections and *Clostridium difficile* colitis rates after 2 years-monitoring in our intensive care unit. Further investigations are needed to precise the impact of dramatic increase in third-generation cephalosporin-resistant Enterobacteriaceae carriage on our results and the best predictors of these hospital-acquired infections.

**Competing Interests**
None.

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**P7**
**Antibiotic therapy against methicillin-resistant staphylococcus aureus is inappropriate in ventilator-associated pneumonia with shock without prior methicillin-resistant staphylococcus aureus colonization**

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**Annals of Intensive Care 2017, 7(Suppl 1):P7**

**Introduction** Ventilator-associated pneumonia (VAP) is the leading cause of nosocomial infections in intensive care units. In case of VAP with shock, most recent international guidelines still recommend an early and broad-spectrum empiric antibiotic therapy focused on any infection due to Gram negative bacilli, including extended-spectrum β-lactamase–producing enterobacteriaceae (ESBL-EB), and MRSA. However, in recent years, there has been a change in the bacterial ecology, particularly a lower MRSA incidence. This study aims to demonstrate that, in patients with VAP and septic shock, an empiric antibiotic therapy with anti-MRSA activity without prior MRSA colonization is inappropriate.
Patients and methods Retrospective cohort study of patients with documented VAP in two intensive care units (ICU). All patients admitted to our ICU’s between January 2010 and December 2014 were included with the following criteria (1) mechanical ventilation for more than 48 h; (2) quantitatively significant culture of lower airways; (3) CPR > 6. At admission and weekly thereafter, in all patients, was routinely performed multidrug-resistant bacterial species colonization by rectal or nasal swab collection.

Results Among 3629 patients under mechanical ventilation hospitalized in our ICU’s, 284 (7.8%) had a confirmed VAP, 172 (60.6%) with septic shock and 112 (39.4%) without. In the septic shock VAP group, 11 patients (6.4%) presented a VAP caused by MRSA. Among them, 10 were colonized by MRSA before VAP occurrence. No infection or colonization by MRSA were found in non septic shock VAP group. Interestingly, among the 284 patients with confirmed VAP, 141 (49.6%) (109 in the septic shock VAP group vs 32 without shock) were treated by anti-MRSA antibiotherapy (vancomycin or linezolid).

Conclusion MRSA colonization screening at admission and weekly thereafter is essential to avoid useless anti MRSA antibiotherapy in VAP patients with septic shock.

Competing interests None.

P8 Systematic screening of multidrug resistant bacteria in a Tunisian ICU

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Introduction Emergence of multidrug resistant (MDR) bacteria is a real problem worldwide and becoming increasingly important. The aim of this study was to describe the frequency and patterns of MDR bacteria following the implement of a systematic screening process at ICU admission in a Tunisian ICU.

Patients and methods Between 1st March 2016 and 31st August 2016, consecutive patients admitted to our ICU following more than 24 h length of stay in another hospital ward, systematic screened for MDR bacteria. Specimens were recovered from nasal, axillary and rectal swabs. Micro-organisms cultures were made on chromogenic media.

Results During the study period 127 patients were admitted to the ICU, 32 of them were transferred from other wards and fulfilled the inclusion criteria. Colonization with MDR bacteria was present in 11 patients (34.3%) (median age 64 [IQR 39–75], 62% males, mechanical ventilation in 59.4%). 72% of patients were referred from the emergency department. During the study period 127 patients were admitted to the ICU, 32 of them were transferred from other wards and fulfilled the inclusion criteria. Colonization with MDR bacteria was present in 11 patients (34.3%) (median age 64 [IQR 39–75], 62% males, mechanical ventilation in 59.4%). 72% of patients were referred from the emergency department. During the study period 127 patients were admitted to the ICU, 32 of them were transferred from other wards and fulfilled the inclusion criteria. Colonization with MDR bacteria was present in 11 patients (34.3%) (median age 64 [IQR 39–75], 62% males, mechanical ventilation in 59.4%). 72% of patients were referred from the emergency department.

Conclusion MDR bacteria (100% extended-spectrum beta-lactamase-producing enterobacteriaceae) is present in more than one-third patients transferred from the emergency department. The risk factors for MDR bacteria acquisition and a strategy to contain this phenomenon are needed in our hospital.

Competing interests None.

P9 Vancomycin-resistant enterococcal faecium bacteremia in an intensive care unit: incidence and risk factors

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Annals of Intensive Care 2017, 7(Suppl 1):P9

Introduction Vancomycin-resistant Enterococcus faecium (VREF) is currently one of the most important etiologies of nosocomial infections in critically ill patients. In the continuum of VREF infections, bacteremia is of special interest, given that overall mortality rates may reach values higher than 60% with an attributable mortality of around 40%. The aim of our study was to determine incidence and risk factors associated with VREF bacteremia in an intensive care unit (ICU).

Patients and methods A retrospective case–control study was performed in the ICU of an university hospital in Tunisia from March 2008 to march 2016. Cases were defined as septic patients with VREF isolated from a blood culture (VREF group). Blood isolates were identified according to standard techniques and Vitek2 (bioMerieux). VREF was defined as an Enterococcus faecium isolate with an MIC of vancomycin > 32 µg/mL by the Etest (bioMerieux) according to the standards of the Clinical and Laboratory Standards Institute (CLSI).

Control patients were randomly drawn from 65 hospitalized patients with vancomycin-susceptible Enterococcus faecium isolated from a blood culture (VSEF group). Medical records of the patients were reviewed. If patients developed several episodes of VREF bacteremia during the study period, only the first episode was investigated.

Results A total of 20 case patients (4.89 per 1000 admissions) and 20 control patients with at least one positive E. faecium blood culture were identified. The demographic and clinical characteristics of the case and control groups were similar, except for mean duration of length of stay, with values being significantly greater in the case group (66 ± 12 vs 24 ± 8, p < 0.001). In 14 cases, VREF bloodstream infections were related to intraabdominal infections; six cases had catheter-related VREF bloodstream infections. Mortality among these bacteremic patients did not differ significantly between those with VREF (14%) and those with VSEF (10%) isolates (p = 0.36). In the univariate analysis, the significant risk factors for VREF bloodstream infections included diabetes mellitus, arterial hypertension, end-stage renal disease, prior exposure to immunosuppressive agents notably corticosteroids, prior receipt of vancomycin before VREF identification and a prolonged length of stay in ICU (Table 10).

Discussion Enterococcus is the third most common cause of nosocomial bloodstream infection. Vancomycin-resistant Enterococcus (VRE) is an important problem in Europe, USA, and Latin America and has been isolated in many other countries. Infections due to VRE have been shown to be associated with significant in-hospital mortality and morbidity. Despite the fact that 80–90% of clinical isolates of enterococci are E. faecalis, most VRE are E. faecium [1]. Similarly, in our

Table 9 Micro-organisms isolated with systematic (MDR) bacteia screening

| Organism                      | N  |
|-------------------------------|----|
| Escherichia coli [n (%)]      | 4  (36.3%)|
| Klebsiella pneumoniae [n (%)] | 4  (36.3%)|
| Klebsiella oxytoca [n (%)]    | 1  (9%)   |
| Citrobacter freundii [n (%)]  | 2  (18.1%)|

ForMDR bacteria acquisition and a strategy to contain this phenomenon are needed in our hospital.
Table 10 Risk factors for VREF by univariate analysis

| Risk Factor          | EFRV (n = 20) | EFRV (n = 20) | p   |
|----------------------|--------------|--------------|-----|
| Diabète              | 15           | 0            | 0.006|
| HTA                  | 15           | 0            | 0.006|
| I Rénale C           | 15           | 5            | 0.03 |
| Corticothérapie      | 17           | 5            | 0.04 |
| Prise de Vanco       | 15           | 2            | 0.048|

institution, the vast majority of VRE bacteremia cases are caused by E. faecium.

**Conclusion** The incidence of VREF bloodstream infections was high compared with literature data.

Several risk factors have been identified and they should be considered in infection control practice to prevent VREF infection or colonization and to reduce their duration. This association has been previously suggested only in studies that have been limited by small numbers of patients and a failure to perform multivariate analysis. The retrospective analysis of a relative small cohort of patients is the major limitation of our study considering that we cannot be certain that we have identified all potential confounding factors.

**Competing interests**

None.

**Reference**

1. Dunn MJ, Breen DP, Davenport RJ, Gary AJ. Early management of adults with an uncomplicated first generalised seizure. Emerg Med J. 2005;22:237–42.

**P10**

Convulsion crisis of the adult in the emergency: epidemiological and clinical characteristics

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**Annals of Intensive Care 2017, 7(Suppl 1):P10**

**Introduction**

The convulsion crisis is a symptom indicating a hyper synchronous discharge of a more or less extensive portion of the cerebral cortex. It is a frequent disease in the emergency (0.3 to 1.2%) [1]. The aim of our study was to report the epidemiological, clinical, therapeutic and evolutionary aspects of the convulsion crisis in the emergency.

**Patients and methods**

Prospective study extending on year (01/01/2015 - 31/12/2015), including all patients aged over 18 consulting in the emergency for convulsive crisis.

**Results**

We included 41 patients. The average age was 45 ± 19 years with a sex ratio of 1.5. The convulsion crisis occurred at a known epileptic in 53.6% of cases. The average duration of crisis was 10 ± 4 min. The average time of consultation was 54 ± 21 min. Generalized tonic-clonic seizures were observed in 30 patients. Partial seizures in 11 patients. The status epilepticus has been reported in 12 patients (29.2%). Discontinuation of antiepileptic therapy was noted in 20 cases (48.7%), Intoxication in 6 cases (14.6%), Hypoglycemia in 5 cases (12.1%), head trauma in 5 cases (12.1%), accident cerebrovascular in 3 cases (7.3%) and preeclampsia in 2 cases (4.8%). The convulsive crisis was accompanied by a post-critical deficit in 6 patients (14.6%) and a head injury in 2 patients (4.8%). A brain imaging was indicated in 18 patients (43.9%). Benzodiazepines were prescribed in 78% of cases and barbiturates in 24.3% of cases. The use of invasive mechanical ventilation was reported in 12 patients (29.2%).

Thirteen patients were hospitalized in intensive care, 12 in neurology and 6 in medicine department. There were no deaths.

**Conclusion**

The management of convulsive crisis in the emergency is multidisciplinary. It is based on a good mastery of the definitions and recommendations on this subject.

**Competing interests**

None.

**Reference**

1. Dunn MJ, Breen DP, Davenport RJ, Gary AJ. Early management of adults with an uncomplicated first generalised seizure. Emerg Med J. 2005;22:237–42.

**P11**

Admission in stroke units (SUs): For which patient if there is only one bed left?

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**Annals of Intensive Care 2017, 7(Suppl 1):P11**

**Introduction**

Stroke is a frequent and serious disease. Stroke units (SUs) have demonstrated efficiency in reducing death and disability after stroke. In France, public health authorities's reports recommended that every patient presenting with stroke symptoms should be admitted in SUs. But beds in these units are still insufficient, so triage seems unavoidable. We aimed to explore admission criteria in SUs in case there is only one bed left. In this situation, is the time window allowed thrombolysis the main criteria for neurologists to accept a patient?

**Patients and methods**

This was a postal and e-mail survey that took place from January 1st to July 31st 2016. An anonymous questionnaire was sent to the 164 neurologists who usually work in the 21 SUs in the region of Ile de France. The survey was in two parts: the first one encompassed questions about the physicians themselves (age, professional status, current job tenure), and, then, in the second part we presented different clinical cases including the type of stroke (ischemic, hemorrhagic, or transient ischemic attack), age ranges of the patients, and different time slots during nyctohemeral period. Finally, data were collected in July 2016.

Results were expressed as absolute numbers and percentages, and for statistical analyses, we used the Fisher’s exact test from the on-line software BiostaTGV.

**Results**

One hundred four questionnaires were completed, representing 63% of all. Majority of physicians were hospital practitioner (53%), and 56% have been working in SUs for more than 5 years. We asked to neurologists if they already refused a patient when it stay one bed in their unit, to keep it for a patient who could be eligible for thrombolysis, and the answer was positive in 64%.

There was no significant difference in accepting or denying regardless of the type of stroke or the different time slot. Concerning the age ranges, we found a significant relationship only with ischemic stroke (p = 0.014), as the oldest patients above 90 years old were more refused than the youngest.

**Discussion**

Our study has several limitations, and probably the main is the risk of declarative biases due to descriptions of practices that differ from those encountered in real life. A more rigorous methodologic analysis, including a prospective study following patients suffering from stroke, would perhaps more reflect the reality.

**Conclusion**

In case there is one bed left in their SUs, neurologists seem to accept most patients, even if thrombolysis is off limits. Only the age ranges may influence their decision if the patient suffers from ischemic stroke. More studies are necessary to confirm or not these results.

**Competing interests**

None.
P12
Assessment of hemorrhagic stroke's management in intensive care unit
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Introduction The management of hemorrhagic stroke (AVCH), considered the second cause of death in the world, has, in recent years major changes. Early diagnosis and optimization of Early and specialized load have also shown their interest on the prognosis of patients
Purpose Through this analytical study, we emphasize the quality of the management of the hemorrhagic stroke.
Materials and methods We realized a retrospective study about 50 patients who suffer from this affection.
Results 50 patients were collected whose average age is 52 plus or minus 15 years male predominance is noted. Hypertension is the most common risk factor for 41% followed by diabetes with a rate of 26%.
Conclusion The outcome was favorable in 14 patients or 28%, and negative in 36 patients who died after surgery.

P13
Optic nerve sheath diameter assessment: it depends of the probe
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Introduction Optic nerve sheath diameter (ONSD) is an easy to obtain parameter, which has been associated to absolute value and variation of intracranial pressure. However, the normal value of this parameter (observed in patients without raised intracranial pressure) remains debatable (from 5 to 5.9 mm) highlighting the importance measurement precision. Is this pilot observational study, we aimed to compare the ONSD values obtained with two different US devices.
Patients and methods During a 2-month period, all consecutive patients recovering from acute disease, orientated, spontaneously breathing, without vasocative support were deemed to be included. Age sex, height, weight, BMI, SAPSII, myopia and other relevant ophthalmologic diseases were recorded. After oral consent, ONSD of both right and left eyes were measured with two distinct US devices: Vivid 7 (GE, equipped with a 8 MHz (4–11 MHz) microconvex probe and a M Turbo, (Sonosite, Bothwell, MA) equipped with a HF38 probe (6–11 MHz). Patients characteristics are expressed as qualitative or quantitative non parametric value as appropriate. The ONSD values obtained with the two devices were compared using Wilcoxon matched-paired test. The bias between the two measurements was expressed according to the Bland–Altman method.
Results During the study period, 25 patients (age: 59 years [50–66], SAPSII: 47[30–64], weight 69 kg [52–86], height 168 cm [158–175]; BMI 23 [20–29], myopia 3 [25]) were studied. Measurements, performed by a senior intensivist skilled in ultrasonography, were always possible with acceptable discomfort. Maximum values observed were 7.9 mm. ONSD values were similar between right and left eyes with the Vivid7 device (5.8 mms 5.7 mm, p = .9) and with MTurbo (4.6 vs 4.7 mm, p = .7) but were significantly greater with the Vivid device (p = .002 and p = .003 for right and left eye respectively). The values were significantly correlated (Spearman coeff r = 0.39 p = .003). No correlation was observed between ONSD values and height, weight or BMI. Myopia was not associated with greater values. The bias (difference mean) 1.16 m provided limits of agreement (±2SD) [−0.0589462, 0.291990] which included all except 3 patients.
Discussion In this small unselected medical ICU patients population with no neurological injury and unlikely to have raised intracranial pressure, the two US devices provided statistically different values with higher values with one of the devices.
Conclusion The ONSD is an easy to obtain parameter but depends on the US devices, which provides significantly different values.

P14
Is there a relevant experimental model of subarachnoid hemorrhage? Systematic review of preclinical literature
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Introduction Delayed cerebral ischemia (DCI) is the main cause of disability after subarachnoid hemorrhage (SAH) but its pathophysiology remains unclear. We currently have no effective treatment to reduce the incidence of DCI. Study of pathophysiology has long suffered from a lack of consensus definition of DCI. It is only recently that experts have proposed that in observational studies and clinical trials aiming to investigate strategies to prevent DCI, the 2 main outcome measures should be: (1) cerebral infarction identified on CT scan or magnetic resonance imaging or proven at autopsy, after exclusion of procedure-related infarctions; and (2) functional outcome [1]. Currently, experimental studies must therefore meet this new definition. The aim of our study is to identify the extent preclinical studies using this new definition of DCI.
Materials and methods Systematic review from PUBMED using key words. Consulting additional bibliographic databases including gray literature, scientific meeting abstracts and review of the bibliography of selected articles completes the research. Inclusion criteria: (1) description and/or modification of an SAH model in rats or mice; (2) study of cerebral vasospasm and/or DCI. This research is performed in accordance with the PRISMA recommendations [2].
Results Seventy-one publications from 47 teams are included. 11 teams are responsible for 77% of publications. 8 different methods are
Table 11 Table of ten most cited studies

| Studies          | Models                   | Vasospasm | Imaging | Histology | Behavioral study |
|------------------|--------------------------|-----------|---------|-----------|------------------|
| Bederson (1995)  | Endovascular perforation | NA        | NA      | NA        | No focal neurological deficit |
| Bederson (1998)  | Endovascular perforation | Yes       | NA      | NA        | NA               |
| Delgado (1985)   | Direct injection         | Yes       | NA      | NA        | NA               |
| Solomon (1985)   | Direct injection         | NA        | NA      | NA        | NA               |
| Prunell (2003)   | Direct injection         | NA        | NA      | Yes       | NA               |
| Jackowski (1990) | Direct injection         | Yes       | NA      | NA        | NA               |
| Sugawara (2008)  | Endovascular perforation | Yes       | NA      | NA        | Yes              |
| Veelken (1995)   | Endovascular perforation | NA        | NA      | Yes       | NA               |
| Gules (2002)     | Direct injection         | Yes       | NA      | NA        | No focal neurological deficit |
| Docti (1986)     | Direct injection         | NA        | NA      | NA        | NA               |

NA not available.

used for induction of SAH. The most used model is the direct blood injection into the cistern magna. Vasospasm is studied by brain imaging (n = 18) mainly in the first 72 h and histology (n = 32) until the tenth day. A positive diagnosis of vasospasm is placed in histology in 28 of 32 studies. Cerebral ischemia is sought in 24 publications including one in cerebral imaging. Neurological deficit is wanted by sensorimotor tests in 13 publications. Three models use animals with comorbidities (hypertension, obesity and chronic inflammation). No model uses female population. Among the 10 most cited studies, none of them is studying cerebral ischemia by imaging.

Discussion Differences with human pathology of SAH are raised about the pathophysiology leading to SAH and its anatomoclinical characteristics. In particular, there is an over-representation of models of posterior circulation. Similarly, study of vasospasm remains the most studied criteria to the detriment of the ischemia and neurological deficit, even though it is the first cause of disability after SAH (Table 11). Furthermore, we can note an over-estimation of the incidence of vasospasm in the included publications compared to the clinical epidemiological data. Study methods of vasospasm and cerebral ischemia may be criticized.

Conclusion Currently we don’t have a murine model of SAH meeting the diagnostic DCI criteria retained in humans. The development of such a model in agreement with the pathophysiological data known to humans remains a relevant objective with the aim of translational research more efficient.

Competing interests None.

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1. Stroke. 2010;41(10):2391–5.
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P16
Brain death in pediatric intensive care: department’s experience
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Introduction Pediatric transplants suffer from organs shortage. Children’s potential organs donors should be enhanced by awareness of the healthcare team and their families, given the socio-economic impact of the transplant. Our study’s aim was to evaluate our potential organs donors from deceased patients in pediatric intensive care during 2015.

Materials and methods We led a retrospective study in pediatric intensive care unit of UHC Ibn Rochd child hospital Abderrahim Elharouchi of Casablanca, over 1 year from 1 January 2015 until 31 December 2015. Were included all patients in clinical brain death. Our study excluded newborns.

Statistical analysis used the epi-info test with significance level P < 0.05.

Results 25 cases were collected, 12% of deaths during 2015, with average age of 7 ± 3, sex ratio was M/W 1.5, the average weight of our population was 34, 72 ± 32 kg, reason for admission was head injury following an PRA for 8 cases, cranial traumatism following defenestration for 2 cases, stroke for 10 cases, and epilepsy for 6 cases. The Glasgow Average on admission was 6.88 ± 3, mean hospital duration was 6.75 days. All patients were ventilated on admission, use of vasoactive drugs were needed for 20 patients.

Brain death diagnosis was made by two intensive anesthetist doctors for each patient looking for brain death clinical signs. Cerebral angiography was made for 10 patients. The procedure of organ donation has been discussed in 17 cases, parents have accepted donation in 2 cases, refusal of both parents in 9 cases, refusal of one parent in 6 cases. In 8 cases the patient died before starting the procedure.

We successfully removed both kidneys and liver for our first donor and just one kidney and cornea for the second one.

Discussion In front of persisting imbalance between the needs and the number of grafts, removal of pediatric organs should be enhanced by optimizing children potential donors and their families management. It is important to encourage the involvement of a health care team and humanly participate with the coordination teams to deal with parents and dare to seek their consent to donate.

Conclusion This study shows out a significant pool of potential organ donors in pediatric intensive care. Hence the need to lead out activities within our society to make of them effective donors.

Competing interests None.

P18
Interest of lung ultrasound in an intensive care unit
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Annals of Intensive Care 2017, 7(Suppl 1):P18

Introduction Interest in bedside lung ultrasound in the intensive care unit has exploded in recent years [1]. The aim of this study was to evaluate the contribution of lung ultrasound in an intensive care unit and its place as a diagnostic tool and therapeutic implication.

Patients and methods Retrospective study extending over 6 months. It included ICU patients who underwent lung ultrasound.
Results Forty patients underwent a lung ultrasound due to various respiratory etiologies: 60% patients for acute respiratory failure, 30% for ultrasound-guided pleural punctures, 20% for exploration of a white lung on chest radiograph. Ultrasonography consists in exploration the chest over the six regions. Of the 24 (60%) acute respiratory failure, the review found: 15 (62.5%) fluid pleural effusions, 2 (8%) pneumothorax, 4 (16%) alveolar-interstitial syndrome and 3 (13.5%) pulmonary condensation. Clinical and gasometric improvement was observed in 70% of patients. Lung ultrasound has allowed a change in management in 43% of patients and has provided new information in 72% of cases and led to successful ultrasound-guided pleural drainage.

Discussion Lung ultrasound has allowed a major improvement of routine practice in intensive care unit. Once the appropriate equipment and training acquired, lung ultrasound has advantages: Safety, speed, acuity, reduced costs and increased patient comfort.

Conclusion Responding immediately to questions for which only sophisticated approaches were used, lung ultrasound untangles these daily problems in the emergency and intensive care.

Competing interests
None.

Reference
1. Rouby JJ. Intensive Care Med.2000;26:1046–56.

P19
How do we set assist control ventilation? Tidal volumes and relations to predicted body weight
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Annals of Intensive Care 2017, 7(Suppl 1):P19

Introduction The use of large tidal volume ventilation can be deleterious and lead to Ventilator Induced Lung Injury. It has been suggested that Lung-protective ventilation using low tidal volume should be used in all critically ill patients. We conducted a study to evaluate the tidal volumes related to body weight prescribed during Assist Control Mechanical Ventilation (ACV) in our Unit and to determine the proportion of patients receiving Low Tidal Volume Ventilation (LTVV).

Patients and methods Retrospective study conducted in a single 12 bed adult intensive care unit (ICU). Patients requiring mechanical ventilation admitted from January to June 2014 were included. Medical files were reviewed to determine the minimal and maximal tidal volumes related to predicted body weight (Vt min and Vt max) delivered during Assist Control Ventilation. LTVV was defined as Vt ≤ 8 ml/kg of predicted body weight (PBW).

Results We included 129 patients (63.5% men) with a median age of 64 years (IQR 52.5–71), a mean SAPS II Score of 46 ± 16 and a median length of mechanical ventilation of 6 days (IQR 2–14.5). Patients were predominantly non-surgical (77.5%) and 18.6% had a diagnosis of ARDS. Mean height was 168.4 ± 9.3 cm, actual and predicted body weight were 72.5 ± 21.7 kg and 63.1 ± 9.8 kg. Median Vt max was 7.9 ml/kg (IQR 7.3–8.9) and 72 patients (56%) had permanent LTVV (Vt max ≤ 8 ml/kg). Median Vt min was 7.4 ml/kg (IQR 6.9–8.2). Women had Vt max and Vt min significantly higher than men [median respectively 9.4 (IQR 8.3–9.8) vs 7.4 ml/kg (IQR 7–8) and 8.4 (IQR 7.5–9.3) vs 7.2 ml/kg (IQR 6.7–7.6); p < 0.0001] (Fig. 11). Only 17% of women had permanent LTVV versus 78% of men.

Discussion Even though the benefit of LTVV applied for all ICU patients is still to be demonstrated, our results show that women are exposed to much more "aggressive" mechanical ventilation with potentially harmful consequences. LTVV seems to be achievable for all intensive care patients since 78% of men admitted in our unit are ventilated with maximal Vt ≤ 8 ml/kg of predicted body weight.

Conclusion A large proportion of patients in our unit are ventilated with LTVV but Mechanical Ventilation could be another example of gender discrimination. Particular attention should be paid when setting tidal volume for women in ICU.

Competing interests
None.

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P20
Respiratory mechanics of passive mechanically ventilated ICU patients
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Annals of Intensive Care 2017, 7(Suppl 1):P20

Introduction Bench test studies are performed to evaluate or compare ventilators. Setting of respiratory mechanics of the lung model should simulate realistic conditions [1]. However, there is no standard for respiratory mechanics properties for normal lung, COPD, and ARDS. Consequently, bench studies use different respiratory mechanics variable which have a big impact on results [1]. This prospective observational study measured respiratory mechanics of passive ventilated ICU patients in order to provide realistic values for bench test studies.

Patients and methods This study was performed in adult mixed ICU of Sainte Musse hospital in Toulon between May 2015 and September 2016. Patients deeply sedated and passively ventilated were included. Exclusion criteria were mixed lung condition, presence of chest tube, prone position, and severe hemodynamic impairment. Patients were classified according to standard definition in: normal lungs, COPD or ARDS. Patients were ventilated in INTELLIVENT-ASV® mode and positioned in semi-recumbent. Respiratory mechanics variables were measured once per patient during the first 24 h after ICU admission together with arterial blood gases. Airway pressure and flow was measured by a proximal pneumotachograph. Static compliance (CSTAT) and inspiratory resistances (RINS) were measured using the least square fitting method [2]. CSTAT was also calculated as the ratio between tidal volume and driving pressure. Expiratory time constant (RCEXP) was measured as the ratio between volume and flow at 75% of maximum expiratory flow and calculated as the product of CSTAT MEAS and RINS. Anova was performed to compare results between each lung condition. Data are presented as median [25th–75th quartiles].
Results Two hundred fifty-seven patients were included: age: 66 [54–74] years; BMI: 25 [22–28] kg/m², SAPS II: 56 [44–66]. The main results are shown in Table 12.

Conclusion This study reports clinically-based values of respiratory mechanics for passive mechanically ventilated ICU patients. These results should help bench studies design to simulate realistic conditions.

Competing interests
None.

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P21
Usefulness of a respiratory comfort scale in titration of the NAVA (Neurally Adjusted Ventilatory Assist) level for patients under spontaneous mechanical ventilation: A physiological study
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Introduction Although Neurally Adjusted Ventilatory Assist (NAVA) is currently used in many intensive care units, the best method for setting the optimal level of assistance in this proportional ventilatory mode is not yet established. In order to determine a method based on aiming the best respiratory comfort of the patient, the authors studied the impact of several levels of assist in NAVA, on the respiratory behaviour and comfort, of patients undergoing weaning of the mechanical ventilation.

Patients and methods This study was single-center prospective, open, randomised, of current care. Patients staying in a critical care unit, undergoing invasive mechanical ventilation for an acute respiratory failure, and recovering spontaneous ventilation, were included. For every patient, an optimal level of assistance in pressure-support ventilation was determined and called Al 100, with which an equivalent level of assistance in NAVA called NAVA 100 was set. From this level, six other NAVA levels were determined, by increase or decrease of 25% stepwise, respectively called NAVA 25, NAVA 50, NAVA 75, NAVA 125, NAVA 150 and NAVA 175, and applied in a randomized order. For every situation, after obtaining a stability of the end-tidal carbon dioxide pressure (ETCO2), pressure, volume, flow, and electromyographic activity of the diaphragm (Edi) were recorded during 5 min. An arterial gazometry was performed, and respiratory comfort was assessed by using the dyspnea numeric scale, the Multi-dimensional Dyspnea Profile (MDP) by Banzett, and the Respiratory Distress Observational Scale (RDOS). All parameters were compared between ventilatory modes, by using variance analysis ANOVA.

Results Ten patients were included. Between NAVA levels, no difference was observed regarding tidal volume (VT, p = 0.143) and arterial carbon dioxide pressure (PaCO₂, p = 0.141). The Ti/Ttotneu ratio and AUC Editot (the area under curve of Edi superior to trigger) raised significantly for NAVA 25, the lowest NAVA level, respectively p = 0.005 and p < 0.001. The dyspnea numeric scale and MDP by Banzett showed no significant difference along the NAVA levels, respectively p = 0.412 and p = 0.107. However, RDOS score was significantly higher for the low levels of assistance NAVA 25 and NAVA 50; p < 0.001.

Conclusion With variation of the NAVA level, our patients showed a respiratory behaviour similar to the findings described previously in the literature. The level of assistance did not seem to influence the patient’s feeling of dyspnea, despite an obvious raising of the ventilatory demand. However, Ti/Ttotneu ratio, AUC Editot and RDOS clinical score appear as potential useful tools to detect under-assistance issues in patients undergoing NAVA.

Competing interests None.

P22
Usefulness of full outline of unresponsiveness score to predict extubation failure in intubated critically-ill patients
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Introduction The aim of the study is to assess the usefulness of the Full Outline of Unresponsiveness (FOUR) score in predicting extubation failure in critically-ill intubated patients admitted with disturbed level of conscious in comparison with the Glasgow Coma Scale (GCS).

Materials and methods All intubated critically-ill patients with disturbed level of consciousness were assessed using both the FOUR score and the GCS. The FOUR score and the GCS were compared regarding their predictive value for successful extubation at 14 days after intubation as a primary outcome measure. The 28-day mortality and the neurological outcome at 3 months were used as secondary outcome measures.

Results Eighty-six patients were included. Median age was 63 [50–77] years. Sex-ratio (M/F) was 1.46. Median GCS was 7 [3–10] while median FOUR score was 8.5 [2.3–11]. A GCS < 7 predicted the extubation failure at 14 days after intubation with a sensitivity of 88.5% and specificity of 68.3% (Youden index = 0.57 95% CI [0.35–0.7]) whereas a FOUR score <10 predicted the same outcome with a sensitivity of 80.8%.

Table 12 See text for description

| Condition       | Normal lung (n = 95) | All ARDS (n = 131) | Mild ARDS (n = 35) | Moderate ARDS (n = 64) | Severe ARDS (n = 32) | COPD (n = 28) | P ANOVA |
|-----------------|---------------------|--------------------|--------------------|------------------------|----------------------|--------------|---------|
| CSTAT MEAS (mL/cmH₂O) | 46 (42–59)          | 34 (29–45)         | 37 (30–46)         | 35 (29–43)             | 33 (29–43)           | 50 (37–70)   | <.001   |
| CSTAT CALC (mL/cmH₂O) | 52 (43–64)          | 40 (32–49)         | 41 (32–51)         | 41 (33–49)             | 38 (30–45)           | 56 (42–72)   | <.001   |
| RINS (cmH₂O s/L)  | 13 (11–15)          | 12 (10–14)         | 11 (9–13)          | 12 (10–15)             | 11 (9–12)            | 22 (16–33)   | <.001   |
| RCEXP MEAS (s)   | 0.06 (0.52–0.72)    | 0.47 (0.41–0.57)   | 0.46 (0.41–0.54)   | 0.47 (0.39–0.57)       | 0.45 (0.38–0.57)     | 1.04 (0.65–1.82) | <.001 |
| RCEXP CALC (s)   | 0.62 (0.51–0.80)    | 0.41 (0.32–0.55)   | 0.44 (0.35–0.57)   | 0.43 (0.30–0.58)       | 0.37 (0.30–0.51)     | 1.17 (0.69–1.70) | <.001 |
and a specificity of 81.7% (Youden index = 0.62 95% CI [0.44–0.77]). The AUC was significantly higher with the FOUR score than with GCS (respectively 0.867 95% CI [0.790–0.944] and 0.832 95% CI [0.741–0.923]; p = 0.014). Both scores had similar accuracy for predicting 28-day mortality and neurological outcome at 3 months.

Conclusion The FOUR score is superior to the GCS for prediction of successful extubation of intubated critically-ill patients.

Competing interests None.

P23 T-tube or pressure support ventilation for the spontaneous breathing trial

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Introduction Spontaneous breathing trial (SBT) is used to test patient’s readiness for separation from the ventilator. The most appropriate method for SBT (T-tube vs pressure support ventilation, PSV) is still a matter of debate. The aim of the present study was to assess SBT and patients outcomes with these two methods in a Tunisian ICU.

Patients and methods This was a cohort study including patients consecutively admitted between 1st July 2014 and 31st August 2016 and ventilated more than 48 h, who underwent their first SBT according to clinical criteria. SBT was performed in our ICU with T-tube before March 2016 and with PSV since then. Clinical characteristics and outcomes were compared between both groups. A multivariate analysis with adjustment on variables significantly different at baseline was made for the prediction of extubation success.

Results During the study period 84 patients under went first SBT (55 patients with T-tube and 29 with PSV). Table 13 summarizes patients characteristics and outcomes.

Multivariate analysis showed no relation between SBT modality and extubation success (extubation without reintubation in the following 48 h) (OR 0.89, 95% CI 0.33–2.38). We also matched 28 patients in PSV group with 28 others from T-tube group (on age, gender and SAPS III), and observed that the rate extubation success was not different (53.6% T-tube vs 57.1% PSV, p = 1.000). ICU mortality was significantly higher among reintubated patients than in successfully extubated patients (75 vs 11.3%, p < 0.001).

Conclusion In our cohort, SBT with PSV or T-tube provided comparable SBT results and clinical outcomes.

Competing interests None.

P25 First-year experience of a French mechanical ventilation weaning centre

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Introduction Economic pressure to maximize resource utilization has resulted in the creation of post-ICU care centers dedicated to patients with weaning failure.

Patients and methods Prospective observational cohort study of patients admitted to a French mechanical ventilation centre located in Essonne; first year experience (January 2015 to December 2015). The 12-bed unit is staffed by nurses with special pulmonary and ventilation expertise, and it features 24-h respiratory therapy supervision and non invasive monitoring (that is ventilator alarms) with signal outputs at each bedside. Pulmonary and critical care specialists are in charge of patients’ treatments and weaning plans. Dietitians and physical therapists are all part of the care team.

Participants 106 patients admitted with a median age of 66.9 years; 60% were male.

70 patients (66%) had tracheostomy before they were transferred in our unit, 54 of them with prolonged mechanical ventilation; 27 patients (25.5%) had non invasive ventilation; 9 patients (8.5%) were dependent on high oxygen flow (>6 l/mm) without mechanical ventilation. All patients came from the ICU.

The duration of ICU stay preceding admission was 236 days. Results The duration of total unit stay was 29 days (6–74 days). The mean IGS score at admission was 29. The mortality rate was 4.7% (3 patients with fibrosis dependent on high oxygen flow, 2 patients—1 with polymetastatic cancer—with tracheostomy and 24 h/ventilator support). 16 patients (15.1%) with worsening conditions returned to the ICU. Among the 85 remaining patients (80.2%), 46 were transferred to a rehabilitation centre, 10 were transferred in acute care hospital in step-down unit, 29 were discharged directly to home. Weaning success was observed in 3 patients with NIV at admission and 24 were educated to self use of NIV.

Of the 16 patients without mechanical ventilation at admission only 3 patients with tracheostomy were discharged with long-term tracheostomy.

Of the 54 patients with invasive mechanical ventilation and tracheostomy at admission:

- 30 were weaned and decannulated
- 7 were discharged with long-term tracheostomy without ventilation
- 14 with long-term tracheostomy were only partially weaned from invasive ventilation
- 1 with tracheostomy and 24/24 ventilator support.

Table 13 Characteristics and outcomes in the two SBT modalities groups before adjustment

| SBT modality          | PSV (n = 29) | T-tube (n = 55) | p    |
|-----------------------|-------------|----------------|------|
| Age (med [IQR])       | 67 [48–74]  | 55 [36–67]     | 0.044|
| Gender (M/F)          | 21/8        | 32/23          | 0.240|
| SAPS III (med [IQR])  | 75 [66.5–80.5] | 67 [60–77]   | 0.021|
| SOFA (med [IQR])      | 8 [7–10]    | 7 [5–10]       | 0.172|
| Reason for mechanical ventilation |               |                |      |
| ARF on CRF (n [%])    | 14 (48.3)   | 16 (29.1)      | 0.101|
| ARF de novo (n [%])   | 8 (27.6)    | 17 (31)        |      |
| Septic shock (n [%])  | 5 (17.2)    | 6 (11)         |      |
| Neurologic (n [%])    | 2 (6.9)     | 16 (29.1)      |      |
| Ventilation duration before 1st SBT (days) med [IQR] | 5 [3–8] | 5 [3–8] | 0.854 |
| SBT outcome           |             |                |      |
| SBT duration, med [IQR] | 60 [30–80] | 60 [30–90]     | 0.489|
| SBT success (n [%])   | 20 (68.9)   | 41 (74.5)      | 0.614|
| Reintubation (n [%])  | 3 (10.3)    | 5 (9.1)        | 0.852|
| ICU LOS (days) (med [IQR]) | 16 [8.5–27.5] | 12 [7–30]     | 0.486|
| ICU mortality (n [%]) | 6 (20.7)    | 10 (18.2)      | 0.778|

Of the patients who were dependent on high oxygen flow (>6 l/mm) at admission 4 were weaned from high oxygen flow at discharge.

Discussion The current data demonstrate that, in our centre, most of the patients with weaning failure can be successfully weaned. Despite a prolonged ICU stay before referral, weaning outcomes were similar to those reported in other countries and organizational models.
Conclusion Patients with weaning failure should be considered for transfer to a specialist weaning centre which can demonstrate favourable short-term outcome.

Competing interests None.

Reference
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P26 COPD patients with acute exacerbation in a Tunisian intensive care unit: What is our practice?
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Annals of Intensive Care 2017, 7(Suppl 1):P26

Introduction Chronic Obstructive Pulmonary Disease (COPD) exacerbations are now the third leading cause of mortality all over the world and in Tunisia. There still are many controversies in their management and mainly the indications of antibiotic treatment and corticotherapy. In Tunisia and despite many valuable studies, only few studies were focused on the "Tunisian" daily practice in case of acute and severe exacerbations of COPD. Thus; we decided to perform this study. We aimed to describe the clinical characteristics of our patients, the blood gas exchange findings, the medical treatment, the ventilator support and to analyze their outcome.

Materials and methods It was a Single-center retrospective study performed in the teaching department of emergency and intensive care medicine in Zaghouran in Tunisia conducted between 1st January 2015 and 31th December 2015. All patients admitted in our ICU for acute exacerbation of COPD were enrolled in the study. Anamnestic characteristics, primary diagnosis, mode of mechanical ventilation were noted. The results of laboratory tests, blood gas data and the outcome of patients were also collected.

Results 40 patients were included in the study. The mean age was 68 ± 11 years with a sex ratio of 4 (32 males and 8 females). The mean SAPS II score was 35 ± 10 with an APACHE II mean score of 22 ± 5. The most common cause of COPD in our patients was tobacco smoke with a mean consumption of 32 ± 10 pack-year history of smoking. The most frequent COPD condition in our patients was chronic bronchitis (n = 29; 72.5%). Six patients received home oxygen therapy. Three patients had lung emphysema and one patient had both restrictive and obstructive syndrome. At admission, the mean Glasgow coma scale score was 12 ± 3. The mean Ph was 7.21 ± 0.13 with a mean bicarbonate level of 32 ± 10 mEq/l. Lung infection was the most common cause of exacerbation (n = 26; 65%). The other recorded causes were: pulmonary edema (n = 3), pulmonary embolism (n = 2) and pleural effusion (n = 5). Twenty-two patients (55%) received systemic corticotherapy and 33 patients received antibiotic therapy. Sixteen patients (40%) did not receive bronchodilator medicines. Fifteen patients (37%) required noninvasive ventilation (NIV). Eighteen patients (45%) was intubated and required invasive mechanical ventilation. Ventilatory support was not necessary in the other seven patients (17.5%). The mean duration of hospital stay in our ICU was 111 ± 98 h. Sixteen patients died in our ICU with a mortality rate of 40%. The most frequent causes of death were septic shock (n = 5), acute respiratory distress syndrome and multiorgan failure (n = 2). Four patients were transferred and twenty patients were discharged from hospital. Only SAPS II was independently associated with poor outcome (p < 0.001).

Conclusion Our patients may be classified severe COPD according to GOLD recommendations. There is a big gap between our daily practice and the international recommendations. We still have a widespread use of corticotherapy and antibiotic therapy. We need effective interventions to implement existing evidence-based guidelines into daily practice in exacerbations of COPD.

Competing interests None.

P27 Serum bicarbonate levels change during ICU stay for hypercapnic COPD exacerbation and impact on long-term survival
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Introduction Acute exacerbation of chronic obstructive pulmonary disease (AECOPD) is linked to morbidity and mortality increase; high bicarbonate levels are linked to chronic hypercapnia and hypoventilation in this population. The consequences of bicarbonate levels change during ICU stay are not well studied. The aim of the present study is to investigate the association between bicarbonate concentration change during ICU stay for AECOPD and long-term survival.

Patients and methods In a prospectively collected database including consecutive patients admitted between 2000 and 2012 for hypercapnic COPD exacerbation in our ICU, we calculated bicarbonate concentration change during ICU stay as follows: ΔHCO3 = [HCO3−] at admission − [HCO3−] at discharge. Patients were split into 3 groups:

- Group 1: Decreased bicarbonate levels: ΔHCO3 ≤ −5 mmol/l,
- Group 2: Small change in bicarbonate levels: −5 < ΔHCO3 < +5 mmol/l,
- Group 3: Increased bicarbonate levels: ΔHCO3 ≥ +5 mmol/l.

Long-term patient’s status (surviving or deceased) was checked by consulting the register of civil status.

Results During the study period 440 patients were consecutively admitted for hypercapnic COPD exacerbation (84.5% males, median age: 68 years, median pH at admission: 7.28 and median HCO3−: 30.6, NIV was the first ventilation modality in 65% and survival at ICU discharge: 84.3%). Blood gas analyses (at ICU admission and discharge), and long-term vital status were available in 239 patients (survival at a median follow-up of 7 years was 31.2%). Bicarbonate levels change were inversely associated with the long-term survival (Table 14; Fig. 12).

Conclusion Our study suggests that bicarbonate levels change during ICU stay for hypercapnic COPD exacerbation could be considered as prognostic factor.

Competing interests None.

P28 Dyspnea in patients with acute respiratory failure requiring non invasive ventilation in the ICU: prevalence, factors and prognosis—a prospective cohort study
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Table 14 Bicarbonate levels change and long-term survival

| ΔHCO3⁻ [med (IQR)] | −5 < ΔHCO3⁻ < +5 (n = 142) | ΔHCO3⁻ ≥ +5 (n = 44) | p       |
|---------------------|-----------------------------|----------------------|---------|
| Long-term survival [n (%)] | 26 (49.1)                  | 57 (40.1)            | 10 (22.7) | 0.027 |

Fig. 12 Kaplan Meier curve, bicarbonate levels change and long-term survival

Table 15 Characteristics and clinical outcomes in patients with and without anemia

|                      | Anemia (n = 133) | Anemia+ (n = 77) | p       |
|----------------------|-------------------|------------------|---------|
| Age [med (IQR)]      | 64 (57–72)        | 71 (65–75)       | <0.001  |
| Male [n (%)]         | 117 (88)          | 67 (87)          | 0.831   |
| Time course of COPD (years) [med (IQR)] | 7 (4–12)       | 8.5 (4.2–15)     | 0.212   |
| pH [med (IQR)]       | 7.30 (7.26–7.34)  | 7.30 (7.27–7.35) | 0.205   |
| HCO3⁻ (µmol/L) [med (IQR)] | 32.8 (28–36) | 31.8 (27.7–35.5) | 0.749   |
| Creat (µmol/L) [med (IQR)] | 83 (66–91)    | 101 (78.2–133.5) | <0.001  |
| GFR (MDRD) [ml/min/1.73 m², med (IQR)] | 83 (70.1–104.5) | 58.1 (42.1–76.1) | <0.001  |
| NIV [n (%)]          | 114 (85.7)        | 67 (87)          | 0.839   |
| Diabetes [n (%)]     | 22 (16.5)         | 15 (15.5)        | 0.580   |
| Hypertension [n (%)] | 33 (24.8)         | 24 (31.2)        | 0.337   |
| Heart failure [n (%)]| 13 (9.8)          | 10 (13)          | 0.497   |
| Ventilation duration (days) [med (IQR)] | 8 (5–11.5)    | 7 (4–10)         | 0.298   |
| LOS (days) [med (IQR)] | 10 (7–14)      | 9 (6–11.5)       | 0.138   |
| ICU mortality [n (%)] | 17 (12.8)         | 8 (10.6)         | 0.665   |

P29
Prevalence of anemia in COPD exacerbations of and impact on short-term prognosis
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Introduction COPD patients have often polycythaemia because of associated hypoxemia especially in patients at the stage of chronic respiratory failure. Little is known about the prevalence of anemia and its impact on prognosis in patients with severe AECOPD admitted to ICU. The aim of this study was to determine the prevalence of anemia in AECOPD patients and its impact on prognosis.

Patients and methods In a prospectively collected database including consecutive patients admitted between 2007 and 2015 for AECOPD in our ICU, we retrospectively assessed hemoglobin levels at ICU admission. Anemia was defined according to WHO criteria: Hb < 13 g/dl in males; Hb < 12 g/dl in females. Continuous variables expressed as median (25–75 percentiles interquartile ranges, IQR) and compared with the Mann–Whitney test.

Results The cohort included 210 patients (median age 67, median pH 7.30, 87.6% males, NIV as first ventilator mode in 86.2%). Anemia was observed in 77 of the 210 patients (36.6%). Median haemoglobin levels were at 10.7 and 14.5 g/dl, in patients with and without anemia, respectively, Table 15 summarizes characteristics and clinical outcomes in the two study groups.
In multivariate analysis anemia was not identified as independent factor associated with ICU mortality. 

**Conclusion** Anemia was observed in one-third of patients admitted to our ICU for severe AECOPD, and was not associated with ICU mortality.

**Competing interests** None.

### P30

**Performance of RV/LV ratio measured at exacerbation in predicting long-term mortality in severe COPD patients**

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**Annals of Intensive Care 2017, 7(Suppl 1):P30**

**Introduction** The identification of specific prognostic factors that predict long-term mortality in COPD patients is paramount for the management of these patients. The objective of this study was to evaluate the performance of the right ventricle (RV)/left ventricle (LV) ratio measured at exacerbation requiring mechanical ventilation in predicting the long-term prognosis in patients with severe COPD.

**Patients and methods** Prospective collection of data on demographics and echocardiography in COPD patients hospitalized in the Intensive Care Unit for hypercapnic decompensation between December 2010 and March 2013. Recorded variables are: age, sex, BMI, medical history, clinical characteristics of the index exacerbation, and blood gas, biological, echocardiographic variables and the mortality at 6 years.

**Results** During the study period 108 were included in this study. The average age was 69 years (range: 43–92 years), 68% were male; the average SAPSII was 27 and average BMI was 26.8 kg/m². The mean PaCO₂: 8.68 ± 1.69 kPa, and pH at admission was: 7.30 ± 0.04. All patients were ventilated with NIV initially. RV was frequently enlarged with NIV initially. RV was frequently enlarged with NIV. The ROC curve depicting the performance of RV/LV in predicting long term mortality is shown in Fig. 13 (AUC: 0.9, a cut-off of 0.7 had the best operative characteristics).

**Conclusion** RV/LV ratio seems a good indicator of long term prognosis in patients with severe COPD.

**Competing interests** None.

### P31

**Serum uric acid (SUA) as a predictor of mortality and future exacerbations of patients hospitalized for severe acute exacerbation of COPD**

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**Annals of Intensive Care 2017, 7(Suppl 1):P31**

**Introduction** Severe acute exacerbation of COPD (AECOPD) reduces quality of life and represents a major burden for health care systems. SUA is well known to increase during hypoxia and systemic inflammation. This final product of purine degradation is one of biomarkers showing promise in the prediction of outcomes in AECOPD [1]. The aim of this study was to assess the possible role of SUA and SUA/creatinine ratio as biomarkers for the prediction of mortality and future exacerbations of patients hospitalized in intensive care unit (ICU) for severe AECOPD.

**Patients and methods** SUA levels were measured in 59 consecutive eligible patients on ICU admission for severe AECOPD between April 2015 and April 2016. Clinical and functional characteristics were compared between patients with levels below and above the median values of SUA and SUA/creatinine ratio. The primary end-point was all-cause mortality at 30 and 90 days. Secondary outcomes included duration of ICU stay, duration of mechanical ventilation and the number of AECOPD in the 6 months period after ICU discharge.

**Results** On ICU admission, age was 69 ± 11 years; SAPS II, 30 ± 9; SOFA score, was 5.0 ± 2.9. Median [IQR] pH was 7.31 [7.28–7.33]; PaCO₂, 57 [53–69] mmHg; PaO₂/FIO₂, 100 [73–203] mmHg; PaCO₂/FIO₂, 239 ± 101. 52.5% had NIV with 45% NIV failure. Overall 71.2% had invasive mechanical ventilation and 59.3% received vasopressors. Mean SUA was 342 ± 90 µmol/L; SUA/creatinine ratio, 4.5 ± 2.1. SUA levels were higher in patients presenting acute heart failure on admission (p = 0.035).

In addition, SUA/creatinine ratio levels were higher in patients having extended evolution time prior to ICU admission (p = 0.05) and those having high SOFA score (p = 0.05).

SUA levels were not associated with increased risk for AECOPD in the 6 months period after ICU discharge.

High SUA was an independent factor associated to circulatory failure in multivariate Cox regression analysis (HR = 12.32, p = 0.037). In multivariate Cox regression analysis, high SUA/creatinine ratio was an independent predictor of 30-day mortality (HR = 15.03, p = 0.032), and it was also independently associated with acute heart failure (HR = 7.08, p = 0.046).

**Discussion** SUA is considered as a useful biomarker in the identification of high-risk patients admitted for mild AECOPD [1–2]. The present study suggests a role of SUA in the prediction of mortality in severe AECOPD associated with heart failure or shock requiring ICU admission.

**Conclusion** High SUA was only independently associated to circulatory failure among patients with severe AECOPD. High SUA/creatinine ratio on ICU admission was associated with acute heart failure and was an independent predictor of 30-day mortality in ICU patients with severe AECOPD.

**Competing interests** None.
P32
Long-term home non-invasive ventilation (NIV) following hypercapnic respiratory failure requiring mechanical ventilation: results of 10 years practice intensive care unit of Monastir
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Annals of Intensive Care 2017, 7(Suppl 1):P32
Introduction
Long-term home NIV is still a matter of debate in patients with obstructive pulmonary disease. Recent studies showed a positive effects on exacerbations frequency and less obvious effects on the survival rate. The objective of this study is to assess the impact of long-term NIV in the long term in patients with persistent hypercapnia at the end of ICU admission for hypercapnic respiratory failure.
Patients and methods
Prospective collection of data in patients hospitalized in the intensive care unit for acute hypercapnic respiratory failure requiring mechanical ventilation between October 2005 and October 2015. Patients with persistent hypercapnia and a diagnosis of COPD, obesity-hypoventilation syndrome, or obstructive sleep apnea had the following recorded data: age, sex, BMI, echocardiographic data, apnea/hypopnea index in the case a polygraphy was performed, long-term NIV duration and observance, the quality of life and the outcome with emphasis on frequency of exacerbations and survival.
Results
During the study period, 110 patients received long-term NIV including 28 COPD, 45 OSA and 37 overlap syndrome. The average age was 65 ± 11 years, BMI: 27. The main comorbidities were: hypertension, diabetes and ischemic heart disease. The baseline pH was 7.38, baseline PaCO2:55 mm Hg. The mean daytime duration of exacerbations was 2.4±1.2 h/day. The mean number of exacerbations per year was 1.24±1.12. The exacerbations were associated with a significant reduction in PaCO2, improved quality of life, and exacerbations’ frequency. Overall mortality at 10 years was 33%.
Conclusion
This study shows the feasibility of NIV in long-term care of patients with persistent hypercapnia following ICU hospitalization for hypercapnic respiratory failure. It also suggests beneficial effects on PaCO2 levels, quality of life and the frequency of exacerbations.
Competing interests
None.

Table 16 See text for description
| Age (ans) | Thrombus+ | Thrombus− | OR (95% CI) | p |
|----------|-----------|-----------|-------------|---|
| 49.9     | 48.6      | –         | 0.757       |

Sex (n, %)

| Male | 23 (25.6) | 67 (74.4) | 1.23 (0.47–3.21) | 0.678 |
| Female | 7 (21.9) | 25 (78.1) |

Neoplasys (n, %) 15 (50.0) 28 (30.1) 2.32 (1.00–5.39) 0.047
Complication (n, %) (slough or sepsis)

| Anticoagulant | 15 (50.0) | 55 (59.1) | 0.69 (0.30–1.58) | 0.379 |
| PAR | 28 (93.3) | 62 (66.7) | 7.00 (1.56–31.31) | 0.004 |
| VAD | 16 (53.3) | 35 (37.6) | 1.89 (0.82–4.35) | 0.129 |

Type of catheter (n, %)

| Double | 5 (16.7) | 47 (50.5) | 1 | 0.001 |
| Triple | 25 (83.3) | 46 (49.5) | 5.11 (1.80–14.49) |

ATB antibiotics, PAR parenteral feeding, VAD Vaso active drugs

Exclusion criteria: Patient admitted with a central venous catheter (CVC), a long term CVC (for dialysis), Presence of thrombo embolic event before catheterization.

All CVC were placed with the help of echoguidance. Each day, we realized a Doppler on the catheter estimating: permeability, Doppler velocity, the presence of thrombus. If this last was detected, we evaluate its length, mobility and adherence to the vein. The entire limb was explored through Doppler echography.

Patients were randomized into 2 groups: with and without thrombus. Data was exploited by SPSS software by univariate then multivariate analysis.

Results
122 catheters were analysed, thrombus incidence was 24.5%.

The comparison between the two groups is mentioned in Table 16. The thrombus was mobile in 83% with a mean length of 4.1 mm.

Conclusion
Thrombo embolic events on CVC are frequent, varying from 8% to 25% of cases and depend on the site. The Doppler helps us to diagnose it quickly and to prevent heavy complications.

In our study, predictive factors were: the neoplasy (p = 0.045), parenteral feeding and the use of triple lumen catheter.

Competing interests
None.

Reference
1. Parienti JJ, et al. Intravascular complications of central venous catheterization by insertion site. N Engl J Med. 2015;373(13):1220–9.

P34
The effects of volume expansion on cardiac output cannot be detected by the changes in heart rate
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Annals of Intensive Care 2017, 7(Suppl 1):P34
Introduction
The primary goal of volume expansion in acute circulatory failure is to increase cardiac output. Nevertheless, techniques that measure cardiac output are not available in every patient. Changes in heart rate could be used instead of cardiac output, since increases in cardiac output induce a decrease in the reflex...
sympathetic stimulation that may be reflected by significant decreases in heart rate. Our goal was to test whether the changes in heart rate could be used to differentiate positive and negative response to volume expansion.

**Materials and methods** We measured heart rate and cardiac output before and after volume expansion (500 mL saline over 10–30 min) performed in patients with acute circulatory failure. We tested the ability of changes in heart rate to detect fluid responsiveness, defined by an increase in cardiac output by more than 15% with volume expansion.

**Results** We analysed 542 volume expansions performed in 242 patients (age 63 ± 14 years old, SAPSII 63 ± 13). The origin of shock was septic in 375 (69.2%) cases, cardiogenic in 24 (4.4%) cases, hypovolemic in 114 (21%) cases, vasoplegic without sepsis in 21 (3.9%) cases and unknown in 8 (1.5%) cases. Cardiac index increased from 3.0 ± 1.1 L/min/m² to 3.7 ± 1.3 L/min/m² (p < 0.01) and heart rate decreased from 96 ± 35 beats/min to 95 ± 31 beats/min (p < 0.01). Cardiac index increased ≥15% in 302 (56%) cases, >30% in 142 (26%) cases and >50% in 61 (11%) cases. Changes in heart rate were not able to detect increases in cardiac index, neither ≥15%, nor ≥30%, nor ≥50% (area under the Receiver Operating Characteristic curves not different from 0.5). Nevertheless, if heart rate decreased ≥20%, which occurred in 15 cases only, an increase in cardiac index ≥30% could be predicted with a sensitivity of 1% (95% confidence interval CI: 0–5%) and a positive predictive value of 25% (3–65%), but with a specificity of 98% (97–99%) and a negative predictive value of 74% (70–78%). Also, if heart rate decreased ≥20%, an increase in cardiac index ≥50% could be predicted with a sensitivity of 2% (0–9%) and a positive predictive value of 1% (1–1%), but with a specificity of 99% (97–99%) and a negative predictive value of 99% (86–92%).

**Conclusion** Except for changes of very large amplitude, decreases in heart rate cannot reliably detect a positive response of cardiac output to volume expansion. Changes in heart rate only allow the detection of large changes in cardiac output.

**Competing interests** None.

**P35**

**Effect of positive end-expiratory pressure and tidal ventilation on the mean systemic filling pressure**

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**Annals of Intensive Care 2017, 7(Suppl 1):P35**

**Introduction**

Mean systemic filling pressure (MSFP) defines the pressure measured in the veno-arterial circulatory system when the cardiac output is nil. Representing the upstream pressure of the venous return gradient, it is considered as a major hemodynamic parameter. Because of the difficulties of its measurement, it is not routinely used in clinical practice but some authors proposed to use heart–lung interaction under mechanical ventilation for its estimation. Our objective was to demonstrate how MSFP is modified by positive pressure ventilation.

**Patients and methods** We conducted a bi-centric non interventional, prospective and observational study to measure the pressure encountered in the circulatory system at the time of death in critically-ill patients on an arterial catheter and a central venous catheter. We included 112 patients with previously inserted catheters and mechanical ventilation with a positive end-expiratory pressure (PEEP) at the time of death. After calibration and supine positioning, arterial and venous pressures were recorded in five conditions: end-expiration and end-inspiration with and without PEEP and finally after disconnection from the ventilator.

**Results** Pressure measured on arterial and central venous catheters did not differ, both representing MSFP. Both arterial and central venous pressures increase with tidal ventilation and with PEEP (Fig. 14). The position of the arterial or the venous central catheters did not have any impact on these results.

**Conclusion** MSFP measured both on arterial and central venous catheters is altered by mechanical ventilation, with a tidal volume and PEEP effect, questioning the use of cardio-pulmonary interactions under mechanical ventilation for its extrapolation.

**Competing interests** None.

**P36**

**Invasive monitoring in intensive units: diagnostic, therapeutic and prognostic benefits**

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**Annals of Intensive Care 2017, 7(Suppl 1):P36**

**Introduction**

Hemodynamic monitoring is an indubitable device in the diagnostic and therapeutic management in critically ill patients. The pulmonary artery catheter (PAC), since 1970, is the gold standard of recommended devices due to the diversity and relevance of provided parameters. Pulse contour cardiac output (PiCCO) represents an interesting alternative to the PAC. However invasiveness and delicacy in the collection and interpretation of data make that the risk/benefit ratio of these techniques is to be worrying. Our purposes were to assess the diagnostic, therapeutic and prognostic utility of invasive monitoring by PAC and PiCCO compared to non-invasive or less common tools.

**Patients and methods** Evaluative retrospective study comparing two arms of ICU patients with invasive versus non-invasive hemodynamic monitoring. Were included, the ICU patients who had hemodynamic and/or respiratory failure and how required the use of hemodynamic exploration. Exclusion criteria were age less than 18 years and the lack of data on the medical record. Outcomes: diagnostic utility was judged on the consistency and quality of data interpretation, therapeutic usefulness on the therapeutic changes and favourable response following these therapeutic adjustments during the 1st 24 h of monitoring, and prognostic assessment was judged on the ventilation period, length of stay and mortality.

**Results** 131 cases of hemodynamic monitoring were included (PAC/PiCCO: n = 71 and common tools: n = 60). The superiority of the invasive monitoring was objectified with the consistency of interpretation of the collected parameters (91.5 vs 78.4%, p = 0.044) and therapeutic changes derived by interpretation (83 vs 66.7%, p = 0.041). No benefit found on the clinical improvement following the modifications guided by monitoring (50 vs 57%, p = 0.72) and (31 vs 47%, p = 0.074) for the hemodynamic and respiratory responses respectively. The duration of ventilation and stay-length did not differ with those of the standard group. A higher mortality was found with age >50 years (OR 1.54, 95% CI [1.05–2.27], p = 0.02) and SAPS II score >35 (OR 1.8, 95% CI [1.09–3.33], p = 0.049).
Conclusion the diagnostic value and therapeutic changes guided by monitoring were significantly in favour of PAC/PICCO technical. No benefits on hemodynamic and respiratory efficiency following these changes with a higher mortality. The invasive techniques keep their interest in intricate situations and where the contribution of non-invasive tools, mainly the echocardiography, is insufficient.

Competing interests
None.

P37
Prognostic value and time course evolution of left ventricular global longitudinal strain in septic shock, preliminary results of a prospective study
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Annals of Intensive Care 2017, 7(Suppl 1):P37

Introduction Septic cardiomyopathy is commonly encountered in patients with septic shock. Most studies suggest no correlation between left ventricular ejection fraction (LVEF) and mortality in patients with septic shock. In the other hand, the initial left ventricle global longitudinal strain (LVGLS) seems to be a better prognostic factor. However its time course evolution remains to be precisely defined.

The present study aims to describe the evolution during the first days of the septic shock and to establish the prognostic value of this novel parameter in critically ill patients.

Materials and methods A prospective observational single center study was performed in the ICU. After approval of the local ethics committee, all patient admitted to the ICU for septic shock without known pre-existing heart disease were eligible. In these preliminary results, 13 of the 100 planned patient were included and analysed.

Echocardiography was performed on the first day, and repeated daily during ICU stay until norepinephrin was stopped. LVEF and LVGLS were evaluated in apical two-chamber, four-chamber and long-axis views. Patients were divided into two groups: survivors and non survivors.

Results Of the first 13 patients included mortality in the ICU was 23%. Left ventricular ejection fraction at admission (LVEF) was not significantly different among the survivors and non survivors (respectively 53% ± 12 vs 28% ± 9; p = 0.089). Initial LVGLS was lower (which indicates better function) in survivors as compared to non survivors (~13% ± 12 vs −6% ± 5; p = 0.044); Time course evolution of LVGLS is displayed in Fig. 15.

Conclusion The preliminary results of the present study suggest that LVGLS has a better prognostic value than LVEF in the initial stage of septic shock.

Competing interests
None.

P38
Prevalence, clinical characteristics and outcomes of left ventricular diastolic dysfunction in the intensive care unit
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Introduction Left ventricular diastolic dysfunction (LVDD) represents about 50% of the causes of heart failure [1]. However little is known about LVDD in the intensive care unit (ICU) setting. The purpose of our study was to determine the prevalence, clinical characteristics and outcome of LVDD in the ICU.

Materials and methods We performed a prospective monocentric study in a medical ICU. Between August 2014 and December 2014, for all consecutively admitted patients, transthoracic echocardiography was realized during the first 24 h after ICU admission. LVDD was established according 2 definitions: 1 complete definition based on American Society of Echocardiography criteria [2], and 1 simplified definition.

Results LVDD prevalence was 61.4% (102/166) according to the complete definition and 56.0% (93/166) according to the simplified definition. LVDD patients were older (66.5 vs 55 years, p < 0.0001). Male sex predominated (67.6 vs 48.4%, p < 0.0001). LVDD was more associated with cardiogenic pulmonary oedema, diuretics use, and new onset of atrial fibrillation, whether the full definition (25.5 vs 0%, p < 0.0001; 52.9 vs 34.4%, p < 0.0001; and 27.5 vs 14.1%, p = 0.044 respectively) or the simplified definition (28 vs 0%, p < 0.0001; 53.8 vs 35.6%, p = 0.002; and 29 vs 13.7%, p = 0.018 respectively). Unlike the full definition, LVDD patients detected with the simplified definition were associated with more important intra-hospital mortality (28 vs 12.5%, p = 0.016).

Discussion LVDD prevalence found in our study is similar as in previous works in the cardiology setting. Higher incidences of atrial fibrillation and cardiogenic pulmonary oedema in the LVDD patients can be explained by a more important left atrial dilation, and a higher incidence of other cardiomyopathies in the LVDD cohort.

Conclusion LVDD is a frequent abnormality in the ICU setting, associated with an important morbi-mortality. LVDD identification should alert the physician for a higher risk of new onset of atrial fibrillation and cardiogenic pulmonary oedema.

Competing interests
None.

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P39

Precision of measurements with transthoracic echocardiography in critically ill patients

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Introduction
We wanted to determine the precision of echocardiographic measurements performed in critically ill patients.

Patients and methods
We included 100 hemodynamically stable patients (age 67 ± 16 years, SAPS II 52 ± 19, 54% mechanically ventilated, 16% with atrial fibrillation and 34% under norepinephrine).

Three successive echocardiography examinations were performed by two different operators with a national echocardiographic diploma, the first and the third by one operator and the second one by the other one. Within each examination, three measurements were performed for each variable without moving the probe from the patient at the end of expiration and averaged.

For every echocardiographic variable, we calculated the precision and the least significant change (LSC), i.e. the minimal change in variable that could be trusted to be significant.

Results
When calculated from the two examinations performed by the same operator, the precision of an echocardiographic examination was 10 ± 9% for velocity time integral (VTI). It is exactly equal to the limit that is usually regarded as acceptable for measures estimating cardiac output. For LV ejection fraction (LVEF), early diastolic peak velocity of the lateral mitral annulus (e’) at Tissue Doppler Imaging and LV end-diastolic area (LVEDA), the precision was 7 ± 9% for VTI. It was 11 ± 12%, 28 ± 19% and 13 ± 9% for VTI. For LVEF, the precision was 7 ± 5%, 8 ± 9%, 17 ± 16% and 10 ± 8%, respectively. In this condition also, LSC of VTI was 14 ± 13%. For LVEF, E, e’ and LVEDA, LSC was 10 ± 8%, 12 ± 12%, 24 ± 23% and 15 ± 11%, respectively. When calculated from the two examinations performed by the two different operators, the precision of an echocardiographic examination was 13 ± 12% for VTI. For LVEF, E, e’ and LVEDA, it was 8 ± 7%, 10 ± 8%, 19 ± 19% and 13 ± 11% for LVEF, E, e’ and LVEDA, respectively. In this condition also, the LSC was 19 ± 17% for VTI. It was 11 ± 10%, 14 ± 12%, 28 ± 27% and 18 ± 15% for LVEF, E, e’ and LVEDA, respectively.

Conclusion
When an echocardiographic examination is performed by averaging three measurements, the precision of echocardiography is almost acceptable for VTI. It is better for LVEDA and LVEF. We are currently investigating the number of measurements that must be averaged within one examination to obtain a satisfactory precision for VTI.

Competing interests
None.

P40

Mixed venous O2 saturation and fluid responsiveness

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Introduction
Hypovolemia is common in intensive care patients. It may reduce cardiac output and O2 delivery relative to tissue need. Low mixed venous O2 saturation (ScvO2) has been used to optimize cardiac output and O2 delivery relative to tissue needs. The use of ScvO2 to predict fluid responsiveness is unclear. Objective: To determine whether ScvO2 is a good indicator of fluid responsiveness.

Patients and methods
We carried out a prospective study in the medical intensive care unit of the teaching hospital in Mahdia over a period of 48 months. All patients with circulatory failure were enrolled. At baseline (t = 0 min), hemodynamic measurements were performed with a PICCO. After baseline measurements and blood sampling, 500 cc of fluids were given over 30 min and hemodynamic measurements were performed at the end of fluid perfusion. Concomitant vasodilator and sedative drugs and ventilatory settings remained unchanged. Fluid responsiveness was defined as an increase in cardiac index >15%.

Results
A total of 68 patients requiring volume expansion were included. The causes of acute circulatory failure were septic shock (n = 45), cardiogenic shock (n = 11), and dehydration (n = 12). Among the 68 included patients, 33 (49%) were responders. ScvO2 was significantly lower in no responders group (62 ± 11 vs 68 ± 12, p = 0.046). However ScvO2 variation was significantly higher in responders group (10 ± 8 vs 6 ± 7, p = 0.041). The area under the ROC curve for ScvO2 was 0.37 (95% CI 0.23–0.5). The best cutoff value of ScvO2 was <55% (sensibility = 36%, specificity = 8 8%, positive predictive value = 75% and negative predictive value = 60%). The area under the ROC curve for ScvO2 variation was 0.65 (95% CI 0.52–0.78). The area under the ROC curve for delta ScvO2 was 0.5 (95% CI 0.43–0.7).

Conclusion
In this study, we can recommend strongly the use of ScvO2 as a predictor of fluid responsiveness.

Competing interests
None.

P42

Association between personality traits and life-saving interventions: a simulation-based study and a questionnaire survey

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Introduction
The application of tactical tourniquet is one of the most important lifesaving intervention (LSI) on the battlefield and in theaters of terrorist attacks. Despite its apparent simplicity, the failure rate of tactical tourniquet application remains high in real life experience. Some personality traits of the combat lifesavers could explain this high failure rate. The aim of this study was to analyze the association between different personality traits and the performance for applying the tactical tourniquet.

Materials and methods
This observational, cross-sectional study concerned French soldiers of two theaters of operations in Africa (SANGARIS in Central African Republic and BARKHANE in Malia), between October 2015 and April 2016. The performance for application of the tactical tourniquet SOFFT® was evaluated during simulation sessions and included the time for application and its effectiveness, demonstrated by the elimination of the popliteal pulse Doppler signal. A performance score included both an application time shorter than 60 s and the interruption of the Doppler signal.

Personality traits were assessed using validated questionnaires: the Self-Esteem Scale of Rosenberg (SES), the HEXACO personality inventory for empathy (HEXACO), the Freiburg Mindfulness Inventory (FMI) and the Büssing Altruism Scale (BAS).

Results
A total of 72 participants (mean age 27 ± 4) were included in the study. The effectiveness rate of tactical tourniquet use was 51%. Only 26% of the participants performed this LSI in accordance with the performance score criteria. Some of the personality traits were associated with significant differences in the performance of the SOFFT® application. Thus, the most empathetic soldiers (high HEXACO levels) had weak performance scores, in comparison with those with a lower HEXACO level (p = 0.016). Participants with a high BAS level were also
less performant for application of the tactical tourniquet. However, combatants with a high SES applied more quickly and more efficiently the tactical tourniquet (Fig. 16).

**Conclusion** Personality traits are associated with significant differences for the performance of tactical tourniquet application, a crucial LSI in setting of both combat and terrorist attacks. Personality traits can be evaluated easily, using validated questionnaires. Finally, the “adaptive learning” could represent an adequate approach to improve the LSI training, according to the personality traits of the trainees.

**Competing interests** None.

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**P43**

**Teaching communication skills through simulation: experience with residents and fellows in paediatric intensive care and anaesthesiology**

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*Annals of Intensive Care 2017, 7(Suppl 1):P43*

**Introduction** Delivering bad news regarding prognosis or care-related damages are difficult situations generating anxiety for patients but also for the medical teams who feel generally insufficiently prepared and trained. We have studied the interest of training paediatric and anaesthesiologist residents and fellows in difficult communication through simulation.

**Materials and methods** This multimodal training with multidisciplinary teachers (psychologists and doctors) included theoretical interactive sessions (behaviour and environment to inform parents in an acute situation, modalities of communication, and knowledge about the defense mechanisms...) and simulation sessions with professional actors playing child’s parents. The scenarios promoted situations after significant events involving a child in PICU or in the operative room. Each scenario was created from real case. Standardized debriefing by teachers and a psychologist was done after each scenario with actors and every participant. A pre and post-test using Likert scale was fulfilled to evaluate the interest of this program and the confidence in this kind of meeting. The results are expressed in median [interquartile] and comparison used the U Man-Whitney test. A p value <0.05 was considered significant.

**Results** Thirty-one medical residents (17 anaesthesiologists, 14 paediatricians) and 5 fellows (1 anaesthesiologist and 4 paediatricians) have participated at this 1 day course training. 44% declared to have previously received a course on communication, and 80% had already experienced, alone, situations with the announcement of bad news.

**PreTest**

- How you consider important the communication with families? 5 [5–5]
- Do you feel trained in communication with the patient or relatives? 2 [2–3]
- Do you feel anxious in situations requiring breaking bad news? 3 [2–3]

**Post Test**

- How do you evaluate this training course? 4 [4–4]
- Does this training course meet your expectations? 4 [4–4]
- Do you feel anxious in situations requiring breaking bad news? 3 [3–3]

97% of participants declared they recommend the training to their colleagues.

**Discussion** The participants reported a significant lack of initial training and a lack in confidence in communication skills with relatives despite their strong interest for this topic. The majority has still encountered such a situation. The post test evaluation showed a significant improvement in confidence even though the level of anxiety remained high after the session. The multimodal aspect of the program combining theoretical courses and realistic simulation with professional actors in a “secured” environment was emphasized by the learners.

**Conclusion** This study confirms the feasibility, the realism and the learners’ satisfaction. It’s of major interest to pursue development of such interprofessional training with others caregivers (nurses...) and other medical and surgical specialists.

**Competing interests** None.
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P44
Usefulness of a sterile in-plane needle-guidance for internal jugular and axillary vein cannulation: a randomized controlled study on an inanimate manikin
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Introduction
International guidelines strongly recommend ultrasound (US) guidance for central venous catheter (CVC) insertion. Compared to anatomical landmark technique, US guidance reduces the number of attempts, and time to successful cannulation of internal jugular (IJ) vein [1]. However, adequate studies to prove this benefit for subclavian/axillary vein catheterization are lacking [2]. This could be explained by anatomical issues but also by technical difficulties. In-plane (IP) technique allows seeing the whole needle’s course but needs to place the needle in the exact middle of the probe in its long axis, and no part of the needle is visualized. In the out-of-plane (OOP) method, only the needle’s tip is visualized, and the axis of the needle is not controlled. Needle guidance systems are aimed at improving needle visualization but data supporting their advantage over conventional US guidance are scarce in the setting aimed at improving needle visualization but data supporting their advantage over conventional US guidance are scarce in the setting of auxiliary vein cannulation. We conducted the present study to compare a sterile in-plane needle-guidance (Infinity Pro; CIVCO Medical Solutions) to standard freehand US guidance during IJ and axillary veins cannulation.

Materials and methods
For both sites, each operator performed venipuncture using three methods: (1) standard OOP (2) standard IP, and (3) IP with tested guide. Procedure was stopped at 180 s when the operator failed to cannulate vein. Punctures were performed on an inanimate manikin (Blue Phantom II, CAE Healthcare St. Louis, MO) using the M-Turbo® device (Sonosite, Bothwell, MA) with a 7.5 MHz linear probe equipped or not with the tested guide. The order of punctures—site (IJ or axillary) and method—was randomized using a 2-by-3 design in a 1:1:1:1:1:1 ratio. The random allocation sequence was generated using a random number table.

The number of attempts (needle passes) before success and time between first skin puncture and successful venous puncture were recorded. Qualitative and quantitative values are expressed as number (percentage), and median (range), and were compared using the Wilcoxon matched pairs test and the Fisher exact test, respectively.

Results
Twenty physicians (median age 22 (22; 27) years, 7 senior physicians, graduated in intensive care medicine, 6 residents, and 3 medicine students) participated to this randomized controlled study. Twelve (60%) declared prior experience in US-guidance for CVC insertion, and none had previously used the tested device. At the axillary site, the tested device significantly reduced the number of attempts compared to both standard techniques (median 1 [1; 1] attempts vs 1 [1; 2.8] for OOP standard, p = 0.021, and vs 1 [1; 2], p = 0.035 for IP standard). Time between first skin puncture and success was significantly reduced with tested guide (median 16.5 [18.9; 39.3] s compared to OOP standard (median 32.5 [9; 180] s, p = 0.029) and tended to be shorter than IP standard method (median 29 [9.5; 96.3] s, p = 0.079). Success rate at first attempt was 85.3% using needle guidance, whereas it was 45% with OOP standard (p = 0.0057) and 55% with IP standard (p = 0.031) methods. No significant difference was observed between needle-guidance and standard methods for IJ cannulation. Compared to IP standard, seventeen (85%) operators preferred in-plane needle guidance device for axillary vein puncture, whereas only 5 (25%) preferred tested device for IJ puncture.

Discussion
These results have yet to be confirmed on a larger sample size and for bedside procedures on patients.

Conclusion
Consistently with previous findings, these results suggest the interest of needle-guidance devices over freehand method for US-guided axillary but not for IJ vein puncture.

Competing interests
None.

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P45
Medical students’ first foray into clinical research through a professional practice assessment: Could it be a win–win experience?
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Introduction
Though efforts have been made in the previous decades to teach medical students the basics of clinical research, their own involvement in such studies infrequently occurs before they become residents or assistants. The medical courses include lessons on experimental and clinical research methods, which may be difficult to integrate out of a concrete context. On the other hand, much has to be learned for medical students in ICU within a short training period. Involving medical students in research could increase their workload, and may affect their evaluation of the training period. We therefore aimed to have medical students making their first foray in clinical research through a practice evaluation. We asked them to evaluate the change in workload and their global satisfaction afterward.

Patients and methods
Four medical students (4th year of medicine) agreed to participate to a clinical trial through an evaluation of medical practices. None of them had had a previous experience in this field. Gastric stress ulcer prophylaxis (SUP) was chosen by the students among other subjects. Two references were provided, in French, to give them clues on how to conduct the research. The first one was a free web access medical thesis exploring the medical practice on ulcer prophylaxis in three ICU in Toulouse. The second was a chapter of a reference textbook. Three weeks thereafter, it was decided in a meeting to have a 2 steps approach. First, they realized an evaluation of the local practices through a daily monitoring of patients admitted during a 2 weeks period. All patients admitted for an estimated period of more than 48 h were eligible. Exclusion criteria were baseline ulcer treatments and active gastrointestinal bleeding. Second, a brief survey of seniors and residents’ knowledge on the subject of matter was performed. In both steps of investigation, the recorded criteria were those described in the French health authority’s recommendations. At the end the study period students participated to statistical analysis and interpretation of results. The study was presented to the department’s staff by the students. A few weeks later, they were asked to evaluate this experience.

Results
During the study period, 27 patients were screened and 14 patients (Mean age 57.4 ± 23.5 years, 4 female, 10 men, mean SAPS II 48 ± 25) were included and observed during a total 105 ICU days (7.5 ± 6.4 days/patient). 6 (42.9%) patients received stress ulcer prophylaxis at least once, and during 71.6 ± 32.6% of their ICU stay. According to French recommendations, SUP was not justified at least once in 5 out of the 6 patients. In 29% of SUP days, no criteria supported this prescription. 8 patients did not receive SUP. These patients were observed during 66 days. According to guidelines, SUP should have been offered in 2 patients during 15 ICU days. In this subgroup of patients, no criteria for SUP was found in only 13 (19.6%) days. Among the studied criteria, a strong link between SUP and enteral nutrition was found (p < 0.0001, Fisher exact test). 8 senior intensivists and 6 residents answered the survey. The estimated prevalence of this
condition was evenly distributed between less than 1% and more 10% among responders. 30% of them ignored whether guidelines were available, and 80% of all responders ignored the strength of evidence of the guidelines. All the known risk factors for stress ulcer were cited, but the two major criteria (mechanical ventilation and coagulation abnormalities) were recognized in less than 30% of responders. 86% of responders declared that only proton pump inhibitor were appropriate for SUP. All students declared that adding this study to the global burden of the training period was manageable. They pointed out that building a study was a difficult task and declared that it was an added-value to the training course, and a good experience in their medical course. They would agree to do it again.

Discussion This study is limited by its single center design, and the little number of participating students and included patients.

Conclusion Helping medical students make their first steps in clinical research seems to be feasible in ICU. The students greatly appreciated this experience and judged it improved their training period. It could also represent a mean for patient management improvement.

Competing interests None.

P46 Does observational role allow skills acquisition in cardiopulmonary resuscitation simulation training? Emilie Duburcq-Gury1, Léa Satre-Buisson1, Thibault Duburcq1, Julien Poissy1, Laurent Robinquet1, Merce Jourdain1
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Introduction High-fidelity simulation is a recognized tool for acquisition and performance improvement in advanced cardiac life support. Our study aims to assess the contribution of attending a high-fidelity simulation session on acquisition of technical and non-technical performances (crisis resource management CRM).

Materials and methods This observational prospective study included second-year post graduate medical residents from Lille2 University, divided into an observational group (A) and an intervention group (B). At the first session, a resident from group A observed a cardiac arrest scenario managed by a resident from group B (B1). A debriefing was performed after each session. The observers and actors expressed feelings, doubts and asked questions to the trainer. Three months later, all residents (A2 and B2) played the role of leader in another cardiac arrest scenario. Their technical performance was rated over 50 points by one trainer and CRM was retrospectively evaluated via a video system by three independent reviewers according to the Ottawa Global Rating Scale (GRS) over 35 points.

Results Thirteen residents from group A and 14 from group B were analyzed. The performances of the group B active residents significantly improved from the first (B1) to the second session (B2), with regard to the technical score (31.6 ± 5.8 vs. 41.1 ± 4.9; p = 0.0004) as well as the Ottawa GRS (18.1 ± 3.2 vs. 22.7 ± 4.6; p = 0.0007). Spectators (A2) had better technical performances than B1 active players (41 ± 3.6 vs. 31.6 ± 5.8; p < 0.0001) and were not different from B2 active players (41.1 ± 4.9 vs. 41 ± 3.6; p = 0.93). There was no significant difference for the Ottawa GRS between A2 spectators and B1 active players (20.4 ± 3.4 vs. 18.1 ± 3.2; p = 0.11), nor between A2 spectators and B2 active players (22.7 ± 4.6 vs. 20.4 ± 3.4; p = 0.17).

Conclusion Our findings suggest that residents who observed a simulation session developed similar technical performances than residents playing for the second time an ACR scenario. This result was not confirmed for non technical performances and further studies are needed.

Competing interests None.

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P47 High fidelity simulation and ECMO in intensive care unit: a multi professional training program to improve skills and self efficacy Thierry Sécheresse1, Mattéo Miquet1, Alexis Simond1, Pascal Usseglio1
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Introduction Technical progress and development of indications may lead non-specialized ICU implanting ECMO, with an investment in equipment and appropriation of new process. Acquisition and retention of new skills to manage these specific situations is essential but difficult because these events are infrequent at an individual level. Full scale high-fidelity simulation conducted in multi-professional team is a way to improve skills and confidence of the teams for these rare situations but with immediate vital risk.

Materials and methods A simulation program on emergency management in relation with ECMO has been jointly set up by members of the intensive care unit for disciplinary expertise in connection with the CEnSIM team for pedagogical expertise. Three priorities have been identified:

- In situ high fidelity simulation
- Multi-professional team simulation (1 intensivist physician, 2 nurses with at least 1 ECMO referrer, 1 caregiver) by 4 h session.
- Program proposed to the whole service to ensure support for uniformity in healthcare

These sessions were conducted in an intensive care room adapted to meet the requirements of high-fidelity simulation. Each session consisted of two simulations each followed by a debriefing. Two objectives were targeted: technical achievement and teamwork.

Beyond the reactions, the main assessment was performed by measuring the evolution of the participant self-efficacy defined by Bandura as "an individual’s belief in its ability to organize and execute the course of action required to produce desired results." Measurements were performed before and after training using analogic visual scale of self-efficacy tailored specifically to each profession.

Results Two test sessions and ten training sessions were conducted over a period of 6 days, allowing the participation of all the physicians and ECMO referral nurses (total participants n = 44: 10 physicians, 4 residents, 20 nurses, 10 caregivers).

This training was appreciated by all participants, both in terms of emotional perception (overall satisfaction: M = 4.9/5; SD = 0.3) and in terms of practical reinvestment (perceived usefulness: M = 4.9/5; SD = 0.3). The results show a beneficial effect of this program on self-efficacy for the three professional categories: Delta (pre-form–post-form) caregivers score = 2.63; p < 0.005; Delta (pre-form–post-form) nurses score = 2.6 nurses; p < 0.001; Delta (pre-form–post-form) physicians score = 3.46; p < 0.001.

Conclusion Inter professional full scale simulation allows the acquisition of experiential knowledge and skills essential to the management of these complex situations. Beyond teamwork and technical skills,
significantly increased self-efficacy is an important factor in achieving optimal behavior in emergencies. In situ simulation performed in interprofessional teams is probably an interesting method to optimize management of ECMO patient. Implementation and perpetuation of such projects need a strong institutional commitment as part of a policy.

Competing interests
None.

P48
Discrepancies in ventilatory settings: frequency, typology and severity in a Tunisian medical ICU
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Introduction
Mechanical ventilation is a life supporting treatment commonly indicated in patient’s ICU [1]. However, discrepancy in ventilatory settings can lead to patient–ventilator asynchrony. This adds a burden on the respiratory system and may increase the morbidity and mortality in the critically ill. The discrepancies in ventilatory settings may be largely underestimated because of a frequent lack of monitoring [2]. The aim of the study was to evaluate frequency, typology and severity of discrepancies in ventilatory settings in a Tunisian medical ICU, and to identify factors associated with patient-ventilator asynchrony.

Patients and methods
An audit observational study was conducted in a 7-beds medical ICU during 1 month period (August 2016). All consecutive ICU patients requiring invasive or non invasive mechanical ventilation were included. The data collected were: patient’s characteristics, initial diagnosis, SAPSII, PaO2/FIO2, ratio, type of mechanical ventilation, ventilatory mode, prescribed ventilatory parameters and interfaces. Waves analyzed, peak inspiratory pressure, plateau pressure, auto-PEEP, volume and inspiratory and expiratory flow waveforms. Patient-ventilator asynchrony was defined as: ineffective inspiratory efforts, auto-triggering, delayed cycling, double triggering and inspiratory waveform distortions.

Results
During the study period, were performed a total of 160 ventilatory settings observations. Mean age was 59.9 ± 16.8 years, mean SAPSII score was 36.3 ± 11.1. 157 (98.1%) patients were on invasive mechanical ventilation and 3(1.9%) were on non-invasive mechanical ventilation. PaO2/FIO2 ratio, the plateau pressure were respectively: 35.5 ± 11.8, 8.4 ± 4.5 and 22.0 ± 4.7 cmH2O.

Discrepancies in ventilatory settings were found in 55(34.4) patients. 23(42) patients had frequent patient-ventilator interactions. Ineffective efforts and double triggering were the two most common asynchronies (43.4 and 13.1% respectively). Patient-ventilator interactions were assessed as severe in 9(16.3). Patient-ventilator asynchronies were associated neither to the severity (SAPS II, PaO2/FIO2) nor to the respiratory system mechanic (peak inspiratory pressure, autoPEEP, plateau pressure and driving pressure). A simple intervention on the ventilatory settings corrected the asynchronies in all the patients.

Conclusion
The discrepancies in ventilatory settings reveals frequent. Patient-ventilator asynchronies are the most observed discrepancies. This could be the consequence of a frequent lack of monitoring.

Competing interests
None.

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P49
The use of the serious game Stayingalive® at school improves basic life support performed by secondary pupils: a randomized controlled study
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Introduction
First aid and cardiopulmonary resuscitation training have received widespread promotion throughout the world. France has promoted a decree establishing the mandatory teaching of basic life support to all French secondary pupils (Décret n° 2006-41 du 11 janvier 2006). Large-scale feasibility of such a training, retraining and assessment remains a big challenge. We conducted a study to evaluate the training efficacy of a serious game designed for cardiopulmonary resuscitation and applied to secondary school students.

Materials and methods
Students of 6th and 7th grades from Paris region were randomized (1:1) to one of the 2 groups using a computerized system. Randomization was stratified by classroom. In the Control group, children received a 30-min teaching on nutritional wellbeing. In the Interventional group, children played a serious game reproducing real-life cardiac arrest situation (3D real-time simulation software, Stayingalive®, iLUMENS-Dassault Systemes) without any adult intervention. Two months later, each student was observed and evaluated by 2 assessors blinded to the study group during a cardiopulmonary resuscitation session using mini manikins MiniAnnePlus® (Laerdal Medical). The assessment scale included 15 items divided into 4 categories (cardiac arrest recognition, call for help, chest compression and use of defibrillator). Primary endpoint was the total score obtained by each student. Secondary endpoints included performance for each item.

Results
A total of 97 children were included in the analysis (Interventional group n = 50, Control group n = 47). Median total score was significantly higher in the Interventional group (7 [6–9] vs 6 [5–8], p = 0.02, Fig. 18). In this group, children performed better on the following items: chest compression rate [10/50 (21%) vs 2/47 (4%), p = 0.01], non-stop chest compression [8/50 (17%) vs 2/47 (4%), p = 0.04], search for a defibrillator [18/50 (38%) vs 3/47 (6%), p = 0.0001], placement of electrodes [44/50 (94%) vs 36/47 (72%), p = 0.005] and defibrillator activation [32/50 (74%) vs 27/47 (54%), p = 0.04].
Conclusion A 30-min experience with a cardiopulmonary resuscitation serious game without any adult intervention improves basic life support performed by secondary school students. It may be further improved by adult supervision, repeated use, and gamification using leaderboards. These results are promising to implement on a larger scale such serious games to improve knowledge and skills of children.

Competing interests
Mini manikins were provided by Laerdal Medical.

PS0 Management of cardiovascular manifestations in patients with Guillain–Barre syndrome associated with Zika virus infection
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Introduccion Although the Zika virus causes predominantly mild symptoms, this mosquito-borne disease has become the newest public health challenge. Meningoencephalitis, myelitis, Guillain-Barré syndrome (GBS) associated with deglutition disorders, arrhythmia, cardiac arrest, myocardial infarction, dysautonomia and microcephaly in new borns are among the most serious recorded complications. The first case of Zika in Martinique, French West Indies, was diagnosed in December 2015. To date, this country of 400 000 inhabitants has recorded over 36,260 suspected cases of Zika virus infection, including 29 cases of GBS with Zika present in urine in 14 of them. Among 519 pregnant women who tested positive, only two cases of microcephaly has been identified. One patient a 76-year-old man died. Our objective was to describe the GBS associated GBS cases complicated by cardiovascular disorders in patients admitted to the Intensive Care Unit (ICU) during the onset of the endemic period.

Patients and methods Prospective study of all GBS-associated GBS in patients admitted in our ICU from 1/12/2015 to 5/25/2016 and presenting at least one organ failure using the SOFA score. During this period, all our ICU patients were systematically tested for dengue fever, Chikungunya and Zika viruses. Cases included in this study were defined by the presence of GBS clinical signs (e.g., distal paresthesias) with biological confirmation using RT-PCR blood and urine and also CSF, and/or positive serology for IgM or IgG.

Results During the study period, 22 Zika-infected patients with GBS were hospitalized in the ICU. The median age was 59 years [19–84] and male/female gender ratio 1.5. The patients presented GBS symptoms (N = 22), acute renal failure (N = 8), cardiac complications (including hemodynamic disorders and arrhythmia; N = 6), bradycardia (N = 6), and ARDS (N = 7), refractory hypoxemia required ECMO VV in one patient. Respiratory failure mainly resulting from swallowing disorders required mechanical ventilation in 20 patients. Patients developed one organ failure (N = 4) and multiple organ failure (N = 18). Four patients presented cardiac arrest. sofa score at 48 h following admission was 10 [2–18]. All the patients received intravenous polyvalent immunoglobulins (0.4 g/kg/day for 5 days). The median duration of ICU stay was 15 days. One patient died.

Conclusion Zika infection may lead to ICU admission due to the development of GBS. Physicians particularly cardiology should be aware of additional cardiovascular complications secondary to GBS-related dysautonomic complications. An emergent need for collaborative and multidisciplinary approach is required to mitigate the imminent Zika outbreak in the Caribbean, by providing diagnostic laboratory facilities, ICU beds availability and assistance for appropriate management of the environment. A prospective study is currently conducted to evaluate the cardiovascular complications associated with fatal cases of Zika infection virus.

Competing interests
None.

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PS1 Mechanisms of sepsis-induced inhibition of malignant tumor growth in cancer mice
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Annals of Intensive Care 2017, 7(Suppl 1):PS1

Introduction The outcome of septic shock in cancer patients has dramatically improved over the last two decades, but the impact on the acute inflammatory insult on the further prognosis of cancer is unclear. Indeed, sepsis-induced immune dysfunctions may directly impact on the prognosis of the underlying malignancy in survivors. We are currently developing a research project in order to investigate the reciprocal relationships between bacterial sepsis and malignant tumor growth. We first reported that sepsis-induced immune suppression promoted malignant tumor growth when tumor cells were inoculated in post-septic mice. Conversely, we observed that sepsis may inhibit the growth of previously established local and metastatic tumors. The present study aimed at investigating the cellular and molecular mechanisms of sepsis-induced tumor inhibition.

Materials and methods We used 8–12 w.o. C57BL/6 J mice. Mice were first subjected to malignant tumor inoculation by subcutaneous injection of the MCA205 fibrosarcoma cell line. Seven days after tumor inoculation, mice were subjected to polymicrobial sepsis induced by cecal ligation and puncture (CLP), without any subsequent antibiotic treatment. Controls were cancer mice subjected to sham surgery. The features of anti-tumoral immune response were compared between CLP- and sham-operated cancer mice at day 1 (early assessment) and day 7 (late assessment) following surgery. The intra-tumoral infiltration of immune cells within tumor tissues and draining lymph nodes was assessed by flow cytometry, and the intratumoral cytokine production was assessed by ELISA. We also investigated the role of bacteria and related pathogen-associated molecular patterns on growth of MCA205 cells. The expression of Tlr2 and Tlr4 genes was assessed by southern blot after DNA amplification.

Results We first confirmed that polymicrobial was able to inhibit malignant tumor growth when applied in mice previously inoculated with fibrosarcoma MCA205 cell line. Not surprisingly, the bacterial load in blood, kidneys and tumors were higher at day 1 in CLP-operated mice than in sham-operated counterparts. Of note, the bacterial contamination of tumors was not sustained at 7 days. We checked that MCA205 cells expressed functional Tlr2 and Tlr4 receptors. MCA205 cells cultured in the presence of Tlr2 and Tlr4 agonists, respectively lipopolysaccharide (LPS) and heat-killed Staphylococcus aureus, did not display accelerated cell growth in vitro, suggesting that the anti-tumoral effect of sepsis is mediated by the septic host's response. The distribution of myeloid and lymphoid immune cells within tumor tissue was quite similar in sham- and CLP-operated mice, with the exception of increased proportions of inflammatory monocytes and Tgd lymphocytes in septic mice. The intra-tumoral cytokine pattern of CLP-operated mice was skewed towards an increased production of both the pro-inflammatory Tnf-a and the anti-inflammatory IL-10 cytokines at day 1.
Conclusion Poly microbial sepsis applied to cancer mice inhibits the local growth of a malignant fibrosarcoma tumor. This anti-tumoral effect is not directly driven by pathogens, but rather involves an enhanced intratumoral inflammatory response. In the light of mild quantitative cell alterations, our results suggest that sepsis induces a functional modulation of immune cells resulting in potent anti-tumoral activity.

Competing interests None.

PS2 Sepsis-like circulatory shock related to haematological malignancies
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Introduction Hematological malignancies may be directly responsible for life-threatening organ failures through tumor lysis syndrome, tissue infiltration, coagulation disorders and obstruction of anatomical structures. Respiratory and renal dysfunctions are commonly encountered in this setting. In addition, some patients may develop a systemic inflammatory response syndrome, responsible for acute circulatory dysfunction, so-called sepsis-like syndrome. Besides advanced life support, the treatment of those cancer-related organ failures relies on timely administration of chemotherapy. In this study, we addressed the features and outcomes of patients with sepsis-like circulatory dysfunction related to hematological malignancies, with particular emphasis on the impact of chemotherapy on organ failures.

Patients and methods This was a 9-year (2007–2016) single-center retrospective observational study performed in a 24-bed medical ICU. Inclusion criteria were age ≥18 years AND presence of a hematological malignancy, either already known at the time of ICU admission or diagnosed during the ICU stay AND development of acute circulatory failure requiring vasopressors without any evidence of underlying infection. Patients with hematological malignancies who received chemotherapy in the ICU were retrieved through the information systems from hospital pharmacy units involved in the delivery of cytostatic drugs, and all medical files were individually checked for inclusion criteria. Data were collected from individual files, and included the overall severity through the APACHE II and SOFA scores computed at the time of ICU admission. The SOFA score was thereafter computed daily. Endpoints were the in-ICU and in-hospital vital status.

Results Over the study period, 24 patients (12 men, 12 females) fulfilled the inclusion criteria. Their mean age was 62 ± 16 years. Most of them we in good functional condition since 22 had a performance status of 0. Two patients were previously immunocompromised (severe combined immune deficiency and kidney transplant). The underlying haematological malignancies were distributed as follows: non-Hodgkin lymphoma (n = 18) including 6 patients with diffuse large B-cell lymphoma, acute myeloid leukemia (n = 4), Hodgkin’s lymphoma (n = 1) and chronic lymphocytic leukemia (n = 1). Seven patients had malignancies newly diagnosed in the ICU. Otherwise, the median time from diagnosis to ICU admission was 8 days (min 2 days; max 4357 days).

The primary reasons for ICU admission were respiratory (n = 11), renal (n = 7), neurological (n = 6) and hemodynamic (n = 5) failures. The mean admission APACHE II and SOFA scores were 18.8 ± 9.0 and 8.6 ± 3.4, respectively. The mean blood lactate level was 4.9 ± 1.4 mmol/L. Mechanisms of organ failures were related to tumor lysis syndrome (n = 10), hemophagocytic lymphohistiocytosis (n = 4), lung (n = 4) and liver (n = 2) malignant infiltration, and disseminated intravascular coagulation (n = 2). Acute circulatory failure requiring vasopressors was present prior to chemotherapy in 18 patients, and was secondarily triggered by chemotherapy in 6 patients. In the first group, administration of chemotherapy was associated with a dramatic improvement in the circulatory conditions (Fig. 19) as assessed by the trend in the hemodynamic SOFA variable. The in-ICU and in-hospital mortality rates were 75 and 79%, respectively. The main cause of death was untractable multiple organ failure (n = 14).

Discussion The mechanism of organ failures in this setting is questionable, but is presumably linked to a massive release of pro-inflammatory cytokines and endogenous danger signals by malignant cells. Modulation of this overwhelming inflammatory response may represent a field of investigation as well as a potential therapeutic target in the future.

Conclusion Urgent administration of chemotherapy is associated with fast improvement in circulatory conditions in patients with sepsis-like shock related to hematological malignancies. However, the overall outcome remains very poor.

Competing interests None.

PS3 PTP1B endothelial gene deletion limits glucotoxicity pathways in endotoxemia model
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Annals of Intensive Care 2017, 7(Suppl 1):PS3

Introduction Glucotoxicity is known to occur during hyperglycem ia and glycemic variability, situations that are associated with poor outcomes in sepsis patients. Although cardiovascular dysfunction is a major cause of mortality during sepsis, the impact of glucotoxicity on the cardiovascular system during sepsis remains unknown [1]. Beneficial effects of protein tyrosine phosphatase 1B (PTP1B) deletion, a negative regulator of insulin signaling, on glucose homeostasis and cardiovascular dysfunction during endotoxemia have been reported. We hypothesized that exogenous glucose administration during inflammation increases cardiovascular dysfunctions by activating glucotoxicity pathways and that this is prevented by endothelial PTP1B gene deletion.

Materials and methods For this purpose, we generated an endotoxin model with glucose administration. EndoPTP1B−/− or wild type (WT) mice received LPS (1 mg/kg) or saline solution followed by five injections of glucose (2 g/kg) or saline solution each hour 12 h after LPS. Endothelial deletion of PTP1B was generated by crossing LoxP-PTP1B with Tie2-Cre mice. The exploration of the cardiac function is performed 20 h after LPS injection (H2O) by the non-invasive
evidences in experimental septic shock.

**Discussion** The absence of variation in the gene and protein expression of CHOP and GRP78 shows a lack of endoplasmic reticulum stress in our model despite cellular stresses generated by glucose.

**Conclusion** In endotoxicin model, the variability of blood glucose aggravates the effects induced by an inflammatory trigger in the cardiac tissue without significant impaired cardiac function in this model. Endothelial deletion in PTP1B involved in the regulation of glucose homeostasis, provides improved glycemic control, with a reduction of pathological activation of iNOS, a marker of abnormal vascular function, and PARP1 marker DNA damage.

**Competing interests** None.

**Reference**

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**P54 Effects of low doses of esmolol on cardiac and vascular function in experimental septic shock**

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**Annals of Intensive Care 2017, 7(Suppl 1):P54**

**Introduction** Administration of a selective β1-blocker, such as Esmolol, in septic shock has demonstrated cardiovascular protective effects related to a down-regulation of inflammation. However, the administered dose systematically induced a reduction in heart rate of approximately 20%, thus limiting its prescription at bedside. The present study aimed to determine whether a non-chronotropic dose of Esmolol still maintains its protective cardiovascular and anti-inflammatory effects in experimental septic shock.

**Materials and methods** Four hours after cecal ligation and puncture (CLP), Wistar male rats were randomly allocated to the following groups (n = 8): CLP, CLP + E1 (Esmolol: 1 mg kg⁻¹ h⁻¹), CLP + E5 (Esmolol: 5 mg kg⁻¹ h⁻¹), CLP + E18 (Esmolol: 18 mg kg⁻¹ h⁻¹). An additional eight rats underwent Sham operation. All rats received a continuous infusion of saline, analgesic and antibiotics 4 h after the surgery. Assessment at 18 h included in vivo cardiac function by echocardiography and ex vivo vasoreactivity by myography. Circulating cytokine levels (IL-6 and IL-10) were measured by ELISA. Cardiac and vascular protein expressions of p-NF-kB/IκB, iNOS, p-AKT/AKT and p-eNOS/eNOS were assessed by Western blotting.

**Results** CLP induced tachycardia, hypotension, cardiac output reduction, hyperlactatemia and vascular hypo-responsiveness to vasopressors. Compared to CLP animals, heart rate was unchanged in CLP + E1 and CLP + E5 but was reduced in CLP + E18. Stroke volume, cardiac output, mean arterial pressure and lactatemia were improved in CLP + E1 and CLP + E5 while vascular responsiveness to Phénylephrine was only improved in CLP + E5 and CLP + E18. Plasma IL-6 levels were decreased in all Esmolol groups. p-NF-kB was decreased in both cardiac and vascular tissues in CLP + E5 and CLP + E18.

**Conclusion** In experimental septic shock, low doses of Esmolol, still improved cardiac function and vasoreactivity. These benefits appear to be associated with a modulation of inflammatory pathways.

**Competing interests**

Dr Kimmoun and Pr Levy received fees from Baxter. The remaining authors have disclosed that they do not have any potential conflicts of interest.

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**P55 Is there an inflammatory rebound upon discontinuation of cisatracurium in critically ill patients?**

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**Annals of Intensive Care 2017, 7(Suppl 1):P55**

**Introduction** For patients treated with cisatracurium, it has been shown that proinflammatory cytokines (IL-1, IL-6, and IL-8) significantly went down during the 48 h of mechanical ventilation as compared to placebo [1]. This anti-inflammatory effect is mediated in part by an inhibition of nicotinic acetylcholine receptor [2]. We did not find in the literature any study exploring the post-discontinuation phase of neuromuscular blocking agent apart the fact that blood oxygenation improved after interruption of cisatracurium infusion. The goal of present study was to demonstrate inflammatory rebound defined from the systemic inflammatory response syndrome (SIRS) criteria after discontinuation of cisatracurium.

**Patients and methods** It was a prospective, single-center, observational study. We included adult patients admitted to the medical intensive care unit of the Croix-Rousse hospital in Lyon, who received mechanical ventilation and cisatracurium infusion between February and August 2016. A rapid intravenous infusion of 15 mg of cisatracurium was administered followed by a continuous infusion of 37.5 mg per hour for at least 24 h. After inclusion in the study, SIRS criteria were monitored daily from cisatracurium onset until the 72th hour after molecule interruption. The SIRS was defined by the presence of at least 2 of the following: temperature >38 °C or <36 °C, heart rate >90/min, respiratory rate >20/min or PaCO₂ <32 mmHg and white blood cells count >12 G/L or <4 G/L or >10% immature neutrophils. Inflammatory rebound was defined as the increase of CHOP and GRP78 after interruption of cisatracurium infusion. The primary outcome was the prevalence of inflammatory rebound 24 h after the cisatracurium discontinuation.

**Results** Thirty-nine patients were enrolled. The prevalence of inflammatory rebound 24 h after the cisatracurium discontinuation was 56.4%. No risk factor in the multivariate analysis was associated with the presence of inflammatory rebound and this rebound did not affect mortality. We also noticed an increased risk of inflammatory rebound with time after cessation of cisatracurium (Fig. 20) and this risk was not
VA-ECLS is associated with serious infectious complications. The objective of this study was to investigate the epidemiology and the risk factors of infection of femoral cannulation site in patients with VA-ECLS after cardiac surgery.

**Results**

At all, 142 patients were investigated. Diagnosis of cannulation infection site was made in 38 (27%) patients. The median time to infection was 10 days [8; 15] (Fig. 21). Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli and Enterobacter cloacae were identified in 16, 12, 12 and 10% respectively. The other bacteria were klebsiella pneumoniae in 15% and staphylococcus in 10%. In univariate analysis, risk factors of cannula site infection were a multiresistant bacterial colonization, a bacteremia event and/or a bleeding event in VA-ECLS cannulation site. In multivariate analysis, only a bleeding event in VA-ECLS cannulation site was identified as a risk factor cannulation site infection. The inhospital mortality was similar in both groups but the hospitalization length stay increased in the «infected» group, 45 vs. 26 days (p < 0.0001) respectively. Furthermore, reoperation increased significantly in the «infected» group, 71 vs. 2% of patients (p < 0.0001) respectively. After reoperation, a reconstructive surgery of the cannulation site was necessary in 45% of cases.

**Conclusion**

In patients treated with VA-ECLS after cardiac surgery, inguinal cannulation site infection occurs in 27% of patients. Bleeding event is the main risk factor.

**Competing interests**

None.

**Reference**

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Fig. 20 Cumulative risk of inflammatory rebound over time after cisatracurium discontinuation

Fig. 21 Delay between the VA-ECLS implantation and infection of the inguinal cannulation site (n = 38)

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**P56**

**Epidemiology and risk factor of femoral cannula infections after venoarterial extracorporeal membrane oxygenation in cardiac surgery**

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**Annals of Intensive Care 2017, 7(Suppl 1):P56**

**Introduction**

The Veno-Arterial Extracorporeal Life Support (VA-ECLS) is an efficient therapy in refractory cardiogenic shock. Nevertheless, VA-ECLS is associated with serious infectious complications. The objective of this study was to investigate the epidemiology and the risk factors of infection of femoral cannulation site in patients with VA-ECLS after cardiac surgery.

**Materials and methods**

We investigated all patients underwent VA-ECLS after cardiac surgery between January 2013 and December 2014, all included in the SARIC database. The infection of inguinal cannulation site was defined as an inflammatory or purulent appearance of the cannulation site associated with a positive quantitative culture of the cannulation site sample. The «infected» and «non-infected» patients were compared by Wilcoxon or Fisher tests. A test of Log Rank and a Cox model was used for univariate and multivariate analysis.
periods we aim to estimate the impact of the amended recommendations issued by European society of cardiology.

**Patients and methods** Retrospective study based on the CUB-REA register, including 31 medical and/or surgical intensive care units between 1997 and 2014. Statistical analysis compared the periods 1997–2009 and 2010–2014.

**Results** We included 4757 definite IE over the two periods, 2848 in 1997–2009 and 1909 in 2010–2014. We observed a significant increase in crude annual incidence (153.0 ± 14.9 vs 217.6 ± 46.4; \(P = 0.018\)) and in incidence density relative to number of stays (0.79 ± 0.08 vs 1.1 ± 0.02%; \(P = 0.004\)). Despite a trend towards increasing number of cases over the first period (1997–2009), slopes of incidence density curves clearly indicate an acceleration of the number of cases since 2009 (\(P = 0.017\)). Patients treated during the second period are significantly older and more severe than those treated before 2009. Surprisingly, use of invasive ventilation, renal replacement therapy, and vasopressor were significantly lower during second period. Contrariwise, resort to surgery has doubled between two periods. ICU mortality is significantly lower in the second period but in-hospital mortality remains unchanged. Concerning pathogens, we found a significant increase in incidence of *Streptococcus* spp. and *Staphylococcus* spp., and no changes concerning intracellular bacteria, *Enterococcus* spp., *Candida* spp. or Gram-negative bacilli.

**Conclusion** Despite some limitations inherent to its retrospective design and to potential diagnostic coding bias, our study highlights a quick shifting landscape in the epidemiology of infectious endocarditis in intensive care, characterized by a strong increase in the incidence and changes in bacterial epidemiology. Restrictive bend in antibiotic prophylaxis guidelines could be substantially responsible for these trends.

**Competing interests** None.

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**PS8**

**Early identification of heparin-induced thrombocytopenia in surgical intensive care patients by using the HIT Expert Probability score: a pilot study**

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**Annals of Intensive Care 2017, 7(Suppl 1):PS8**

**Introduction** Heparin is widely prescribed in patients admitted in surgical intensive care units (SICU) to prevent venous thromboembolic events. Heparin-induced thrombocytopenia (HIT) is a rare but potentially life-threatening complication of heparin therapy. HIT requires the emergent discontinuation of heparin and the prescription of an alternative anticoagulant therapy that could be difficult to manage in SICU patients and enhance the risk of hemorrhagic complication. The early diagnosis of HIT in SICU patients remains a challenge. As thrombocytopenia could reveal several SICU complications, the 4T score of Warkentin is a useless tool to efficiently discriminate patients having or not HIT and the biological confirmation of HIT is delayed. The HIT Expert Probability (HEP) score has been reported to have a higher predictive value than the 4T score in non-ICU patients. The purpose of the study was to compare the HEP score to the 4T score in the early diagnosis of HIT in SICU patients.

**Materials and methods** We conducted a one-center prospective observational cohort pilot study (www.ClinicalTrials.gov: Identifier: NCT02790567), included all consecutive patients admitted in our SICU between October 2013 and May 2015 and suspected to have HIT. Non-inclusion criteria were pregnancy, age <18 years old and treatment with fondaparinux. The day the diagnosis of HIT was suspected, the HEP and the 4T scores were calculated and the following blood analyses were performed: the ID-PaGIA Heparin/PF4 test, the ELISA test, the heparin-induced-platelet-aggregation (HIPA), and the serotonin release assay (SRA). After completion of the study, all medical files were reviewed by a multi-disciplinary independent committee to discriminate patient having (HIT group) or not (SAFE group) a HIT. The final diagnosis was based on the medical history of the patient, on the time-variation of the platelet count while heparin was discontinued or not, and on the results of the venous Doppler of the 4 limbs. The committee was blinded from the value of the HEP and of the 4T scores, and from the results of the ID-PaGIA Heparin/PF4 test. The ROC curves of the HEP and of the 4T score were constructed. The specificity (Sp), the positive predictive value (PPV), and the negative predictive value (NPV) of the HEP and of the 4T scores were calculated, and the areas under the ROC curves (AUC) were compared by using a Chi2 test. Data are presented as median [interquartile range] and number of patients (percentage).

**Results** From the 119 patients included, 6 (5) patients had a HIT (age: 66 [59–74] vs 70 [54–80] year, p = 0.62; male: 80 (71) vs 6 (100), p = 0.18; IGSII score value: 54 [46–68] vs 46 [38–62], p = 0.25; respectively in the SAFE and in the HIT groups). The global incidence of HIT during the study was 0.43%. The ROC curves are presented in Fig. 22. The Se, the Sp, the PPV and the NPV of a HEP score ≥5 were respectively 100 and 50%, 93 and 86%, 50 and 22%, and 100 and 96%. All patients in the HIT group had a HEP score ≥5. The AUC was significantly higher for the HEP score (AUC [95% confidence interval] 0.967 [0.922–1.000]) than for the 4T score (AUC [95% CI] 0.707 [0.449–0.965]) (p = 0.035).

**Conclusion** Based on a small prospective observational pilot study, a HEP score ≥5 could have a higher predictive value than a 4T score ≥6 in the early diagnosis of HIT in SICU patients. This result needs to be confirmed in a larger multicenter study.

**Competing interests** None.
P59 Purifying efficiency of CVVHDF and MARS during a simulated intoxication pentobarbital
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Annals of Intensive Care 2017, 7(Suppl 1):P59

Introduction Drug poisoning is a frequent cause of hospital admission especially in intensive care unit (ICU). Despite advances in treatment, hospital mortality of severe acute poisoning admitted in ICU seems to increase. Purifying methods, continuous haemodiafiltration in venovenous (CVVHDF) and molecular adsorbent recirculating system (MARS) were developed with promising clinical results [1]. However, no analytical study has quantified their accurately purifying efficiency. It has not been assessed efficiency of the different compartments of MARS nor its advantages over other methods of dialysis and filtration. The objective of this study was to quantify the purifying efficiency of the different compartments of the CVVHDF and MARS and to compare their respective efficiency in an ex vivo model in the most favourable conditions for these methods to assess their maximum capacity purification.

Materials and methods We performed an ex vivo study based on a manipulation bench simulating intoxication pentobarbital at a plasma concentration of 40 mg/l injected into a central compartment (5 l) devoid of transporter proteins (200 mg of pentobarbital). The EC extraction coefficient [EC = (in concentration – out concentration)/in concentration] were calculated for each compartment of CVVHDF and MARS as well as the amounts withdrawn by the sum each compartment allows to assess the overall capacity of each technique.

Results At the end of 6 h simulation with CVVHDF, the remaining material in the central compartment was 3% of the total quantity injected. The cumulative amount removed in total effluent was equal to 95%. The non-recovered amount was equal to 2%. EC CVVHDF was almost constant with an average value of 14% with large variations however (range 2.5–27%). It was not observed to release at the end of manipulation. At the end of a 6 h simulation with MARS, the remaining amount in the central compartment was undetectable. The cumulative amount removed in total effluent from the hemodiafiltration column was equal to 11.5% the amount removed by carbon column was 88.5%. The sum of the amounts removed by the effluent and the carbon column reflects the total amount injected. The EC hemodiafiltration and charcoal hemoperfusion were 25 and 70% respectively. Purification of pentobarbital was complete after 3 h of a MARS session.

Discussion MARS is the most effective purifying method in a pentobarbital-simulated intoxication. This rapid and complete treatment is mainly due to purifying coal capacity. It was not observed the cartridge saturation for the amount of administered pentobarbital.

Conclusion Future studies should determine the parameters that may affect the treatment capacity by both methods.

Competing interests None.

P60 Plant poisoning: still a current intoxication
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Annals of Intensive Care 2017, 7(Suppl 1):P60

Introduction Plant poisoning is generally not life-threatening. Its occurrence in children is commonly accidental. In adults it results from a suicide attempt or from its use for addictive or therapeutic purposes. Because of their specificity in clinical presentation and outcomes, we have conducted this study.

The objective of this study was to report all plant poisoning cases collected in Tunisian toxicological intensive care unit, with their epidemiological, clinical characteristics and outcomes.

Patients and methods A retrospective study was performed between January 2007 and December 2015. Epidemiological data and clinical outcomes were reviewed. Data were analyzed using SPSS.

Results During the study period, 38 patients were included. Sex ratio was 1. Mean age was 43 years (5–65). Poisoning was accidental in 88% of cases. Most frequently incriminated plants were Datura stramonium (40%) with an anticholinergic toxidrom present in all cases, Ricinus communis (23%) with gastrointestinal manifestations (present in 88% of cases), Nerium oleander (9%) with digitalis toxicity-like symptom in 66% of cases, Hyoscyamus Niger (12%) with anticholinergic symptoms in 75% of cases and hallucinogenic effects in all cases, Atractylis gummifera (9%) with gastro-intestinal and acute liver failure symptoms, and Peganum harmala (9%) with only gastro-intestinal effects. All patients received supportive care. Mortality rate was 8, 5%, interesting children, and was secondary to multi-system organ failure due to ingestion of Atractylis gummifera.

Conclusion Each year 2.2% of our poison center calls report exposures to toxic plants. Most of these exposures are of minimal toxicity largely because of the fact that they involve pediatric ingestions, which are of low quantity. Public education is important to minimize these poisonings and must be oriented primarily towards children to reduce mortality.

Competing interests None.

P61 Toxic effects of rhamnus alaternus: a collective intoxication
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Annals of Intensive Care 2017, 7(Suppl 1):P61

Introduction Herbal remedies have been used for centuries to treat a variety of diseases. Mediterranean Buckthorn (Rhamnus alaternus) has been used for therapeutic purposes and no toxicity effects have been documented. Rhamnus alaternus (Rhamnaceae) is a small tree located mainly in the North of Tunisia, where it is known as “Oud El-khir”. It has traditionally been used as a diuretic, laxative, hypotensive drug and for the treatment of diabetes, hepatic and dermatologic complications. Previous phytochemical studies have shown potent antioxidant, free radical scavenging, antiinflammatory and antigenotoxic activities of flavonoids and phenol isolated from Rhamnus alaternus roots and leaves.

Patients and methods It was a retrospective study reporting a family collective poisoning which occurred in the region of Zaghouan in Tunisia in July 2015. All the members of the family ingested accidentally a traditional preparation of a plant “Oud El-khir” in a juice prepared for a traditional marriage.

Results On 1st July 2015, a family composed of seventeen members (ten men and seven women) was admitted to the teaching emergency and intensive care department of the regional hospital of Zaghouan (Tunisia). All members of the family presented dizziness, weakness, anorexia and dyspnea. They reported the ingestion 10 h before of a juice in a traditional marriage. This juice was prepared using a plant...
called “Oud El-Khir” because of its capacity to be a lucky charm for the new married. The mean blood pressure was 130/60 mmHg. All members of the family experienced nausea, vomiting, anuria and hematuria. On physical examination, five members of the family had myalgia without other clinical signs.

For all patients, urine reports and sputum smear were negative (three times) for pulmonary tuberculosis. Hepatitis B and C serology were also negative. Chest X-ray was normal; Blood and urine culture were negative. In renal ultrasonography performed in five members of the family, there was a significant difference in kidney sizes and the corticomedullary differentiation was altered. Laboratory tests showed hyperglycemia and renal failure with metabolic acidosis in ten patients. Three dialysis sessions were performed. Samples of the herbal decoction were obtained from the juice. It was a dark brown suspension with fine brown deposit and a clear supernatant. It smelled a strong penetrating odor. Samples of both Rhamnus alaternus roots and its decoction were sent to be analyzed in the laboratory of toxicology in the Center for Emergency Medical Assistance of Tunis in Tunisia.

Screening by GC–MS of both Rhamnus alaternus roots and infusion extracts, revealed the presence of anthraquinone glycosides such as 4,5-dihydroxy-9,10-dioxaanthracene-2-carboxylic acid (Rhein), 1,8-dihydroxy-3-(hydroxymethyl)-9,10-anthracenedione (Aloe-emodin) and 1,8-dihydroxy-3-methoxy-6-methylanthracene-9,10-dione (Physcion). The retention times were 8.95, 9.67 and 10.25 min respectively. Anthraquinone glycosides were detected in a dichloromethane extract and ethyl acetate extract at pH value = 9 and only in a dichloromethane extract at pH value = 7 by GC–MS analysis.

**Conclusion** Rhamnus alaternus can be toxic when used in an abusive way besides its strong antibacterial, antioxidants and anti-diabetic activities. To our knowledge, this is the first report of cases of renal failure and rhabdomyolysis which is possibly associated with an accidental consumption of Rhamnus alaternus roots. We present these cases to illustrate the role of both clinical and biological investigations in handling cases of herbal poisonings. We aimed also to increase awareness among emergency physicians about patients presenting to the emergency department with unexplained symptoms (renal failure, rhabdomyolysis...) that requires prompt diagnosis so that such life-threatening complications can be avoided.

**Competing interests** None.

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**P62** Electroencephalographic patterns of lithium poisoning: a study of the effect/concentration relationships in the rat

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**Annals of Intensive Care 2017, 7(Suppl 1):P62**

**Introduction** Lithium overdose may result in encephalopathy and electroencephalographic abnormalities. Three poisoning patterns have been identified based on the ingested dose, previous treatment duration and renal function. Whether severity of lithium-induced encephalopathy depends on the poisoning pattern is not established.

**Materials and methods** We designed a rat study to investigate lithium-induced encephalopathy and correlate its severity to plasma, erythrocyte, cerebrospinal fluid and brain lithium concentrations previously determined in rat models mimicking human poisoning patterns. Lithium-induced encephalopathy was assessed and scored using continuous electroencephalography.

**Results** and **Conclusion** that lithium overdose was consistently responsible for encephalopathy which severity depended on the poisoning pattern. Acutely poisoned rats developed rapid-onset encephalopathy which reached a maximal grade of 3/5 at 6 h and disappeared at 24 h post-injection. Acute-on-chronically poisoned rats developed persistent and slightly fluctuating encephalopathy which reached a maximal grade of 3/5. Chronically poisoned rats developed rapid-onset but gradually increasing life-threatening encephalopathy which reached a maximal grade of 4/5. None of the acutely, 20% of the acute-on-chronically and 57% of the chronically lithium-poisoned rats developed seizures. The relationships between encephalopathy severity and lithium concentrations fitted a sigmoidal Emax model based on cerebrospinal fluid concentrations in the acute poisoning and brain concentrations in the acute-on-chronic poisoning. In the chronic poisoning, encephalopathy worsening paralleled the increase in plasma lithium concentrations.

**Conclusion** Severity of lithium-induced encephalopathy is dependent on the poisoning pattern, previously shown to determine the lithium amount accumulated in the brain. Our data supports that electroencephalography is a sensitive tool to score lithium-related neurotoxicity.

**Competing interests** None.

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**P63** Electrocardiographic changes in amitriptyline poisoning

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**Annals of Intensive Care 2017, 7(Suppl 1):P63**

**Introduction** Amitriptyline is one of the common drug poisonings that induces severe cardiovascular and neurological complications. The incidence of electrocardiogram changes in Amitriptyline poisoning was not explored enough. Therefore, we conducted this study to determine the incidence of electrocardiogram abnormalities in Amitriptyline poisoning.

**Patients and methods** It was a retrospective study conducted from January 2012 to July 2016 in a toxicological unit including all admitted patients for Amitriptyline poisoning. A 12-lead electrocardiogram was carried out and analyzed at admission and every 6 h.

**Results** One hundred and fifty patients aged 29 ± 12 years were included; their sex ratio was of 0.19. Among them, only 6 had a history of cardiovascular disease. A loss of consciousness was noted in 58% of patients (n = 87) and electrocardiogram changes in 42% of patients (n = 63). The most common electrocardiogram changes were sinus tachycardia (108 ± 13 beats/min) (60%), widening of the QRS complex (16%) and right bundle branch block (12%). A change in the S-T was observed in 4 patients and dominant secondary R wave (R > 3 mm) in aVR in 3. Forty-four percent of patients with electrocardiogram changes were comatose. No patient had developed hemodynamic instability especially in the case of widening of the QRS. These electrocardiogram abnormalities took 6 to 36 h to regress without specific treatment.

**Conclusion** Although electrocardiogram changes are common in Amitriptyline poisoning particularly in a patients with loss of consciousness, it rarely induced hemodynamic instability.

**Competing interests** None.

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**P64** Tramadol-related neurotoxicity in the rat: contributions of the different neuromediators and effects of potential antidotes

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**Annals of Intensive Care 2017, 7(Suppl 1):P64**

**Introduction** Tramadol, an opioid analgesics used to treat moderate to severe pain, is responsible in overdose for coma, respiratory depression, seizures and serotonin syndrome. The exact role of...
naloxone to reverse tramadol-related effects is debated. We aimed at investigating the pathways involved in tramadol-related neurotoxicity and seizures in the rat, by using various antagonists of the different tramadol-mediated effects including naloxone, cyproheptadine, feoxofenadine and diazepam and determining the turnover of brain monoamines.

Materials and methods Body temperature (using telemetry), respiratory effects (using plethysmography) and neurological effects (using clinical scales and EEG) were studied. Brain (frontal lobes) monoamines (serotonin, dopamine and norepinephrine) and their respective metabolites were measured using HPLC coupled to fluorimetry. For each animal and each time, we calculated the difference between the parameter value at that time and baseline and the area under the curve of its time course. Comparisons were performed using two-way ANOVA followed by post-tests using Bonferroni correction.

Results Tramadol induced sedation (p < 0.01), seizures (early onset and peaking at 30 min) and increase in inspiratory time (p < 0.001) as well as a non-significant trend to hypothermia. Diazepam completely suppressed seizures Naloxone prevented tramadol-related sedation and respiratory effects while did not inhibit seizures. In contrast to cyproheptadine which exhibited no effects, feoxofenadine partially reduced seizures, suggesting the involvement of a histaminergic pathway. Monoamines turnovers were significantly reduced in the presence of diazepam (p < 0.01), suggesting that diazepam-mediated prevention of tramadol-induced seizures could be related to the inhibition of monoamines metabolism in addition to its usual GABAergic effects.

Conclusion Tramadol-induced sedation and respiratory effects are mediated by mu-opioid receptors. Seizures involve complex mechanisms including histaminergic but not serotoninergic pathways. Diazepam-related anticonvulsive activity to prevent tramadol-induced seizures may be related to the inhibition of monoamines metabolism in addition to its GABAergic effects.

Competing interests None.

P66

Tramadol poisoning in the intensive care unit: clinical presentation and prognostic value of plasma tramadol concentration of on admission

Introduction Tramadol poisonings are significantly increasing due to the rise in prescriptions since dextropropoxyphene banning from the European market in 2011. Tramadol-related analgesic effects are mediated by its antagonist activity on the norepinephrine and serotonin transporters in addition to the agonist activity of its major active metabolite M1 on the mu-opioid receptors. Thus, tramadol overdose may result in various toxicities including central nervous system depression, seizures and serotonin syndrome. The relative prevalence of each of these complications is debated. We aimed (1) to describe the clinical features in tramadol-poisoned patients and (2) to study the prognostic value of the plasma concentration of tramadol and its metabolites on ICU admission.

Patients and methods We conducted a prospective single centre observational study including all tramadol-poisoned patients admitted to the intensive care unit (ICU) from 2012 to 2016. The plasma concentrations of tramadol and its metabolites were determined using high-performance liquid chromatography coupled to mass spectrometry. Subgroup comparisons were performed using Chi-2 and Mann–Whitney tests.

Results Forty-two tramadol-poisoned patients (41 years [26; 55], median [25; 75 percentiles], 30 females et 12 males; 90% with poly-intoxications; presumed ingested dose: 2000 mg [1000; 4000]; plasma tramadol concentration on admission: 1.48 mg/L [1.17; 2.34]) were included in the study. The patients presented consciousness impairment (Glasgow coma score: 13 [6; 15]), opioid syndrome (48%), serotonin syndrome (36%) and seizures (24%). Life-threatening complications occurred including pre-hospital cardiac arrest (10%), cardiovascular failure (31%), aspiration pneumonia (34%), disseminated intravascular coagulation (5%) and fatality (7%). There was a significant relationship between plasma tramadol concentration measured on admission and the risk of seizure onset (p < 0.05). Patients presenting opioid syndrome on admission significantly had lower plasma tramadol concentrations on admission than patients presenting serotonin syndrome (p < 0.03).

Conclusion Tramadol poisoning may result in significant morbidities requiring invasive ICU management. Onset of seizures is not related to the serotonin syndrome. Measurement of the concentrations of tramadol and its metabolites on ICU admission seems helpful to predict the kind of toxic syndromes presented and the nature of further complications.

Competing interests None.

P67

Purpura fulminans mortality factors in pediatric intensive care department: about 28 cases

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**Introduction** Purpura fulminans (PF) is one of the biggest pediatrics emergencies. It is a septic shock associated with an extensive purpura complicating most often meningococcal septicemia. With a non casual mortality despite of advances in pediatric intensive care. Our study’s aim is to study purpura fulminans mortality factors in our pediatric intensive care unit.

**Materials and methods** We led a prospective study of a series of 28 cases of purpura fulminans collected in our pediatric intensive care unit, Abderrahim Harouchi child hospital of Casablanca, over a period of 2 years from January 2014 to December 2015.

Statistical analysis used the epi-info test with significance level P < 0.05.

**Results** Our series analysis underlined a slight male predominance (53.6%) and incidence peak in patients of less than 4 years (67.8%). Most of the patients were from a low socioeconomic level (65%). And earlier evolution period to hospitalization relatively long between 4 h and 7 days was noted in our series.

The main symptoms were fever (100%), purpura (28.6%), digestive disorders (42.8%) and neurological signs mainly consciousness disorders (28.6%). The shock, infectious syndrome, purpura, neurological and respiratory disorders dominated the clinical background at admission. Hyperleukocytosis to polynuclear neutrophils was found in 67.9% of the cases, C-reactive protein was positive in 100% and the hemostatic assessment underlined disseminated intravascular coagulation in 78.9% of the cases. The causative organism was isolated only in 14 cases: meningococcus B in 11 cases (78.6%), pneumococcus in one case, Haemophilus in one case and Acinetobacter in one case.

Therapeutic schema used was based on fluid replacement and antibiotic therapy for all patients, vasoactive drugs (50%), corticosteroids (100%), mechanical ventilation (46.4%) and symptomatic treatment.

The evolution was marked by death in 42.9% of the cases, two patients showed out complications of skin necrosis and distal ischemia requiring necrosectomies. Recovery without sequelae was achieved in 50% of the survivors.

**Conclusion** Mortalities factors found in our series was the time between the first symptoms and hospitalization exceeding 48 h, presence of extensive purpura of a shock on admission, presence of seizures, the use of mechanical ventilation and the presence of disseminated intra vascular coagulation.

**Competing interests** None.

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**P69 Predictors of mortality during purpura fulminans in a developing country**

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**Annals of Intensive Care 2017, 7(Suppl 1):P69**

**Introduction** Purpura fulminans (PF) is a major cause of mortality and morbidity in children. Despite therapeutic advances in the management of PF, mortality remains high up rates that exceed 50%. In Tunisia, the profile of such serious disease has not yet been described. The aim of this study was to determine the predictors of mortality in patients admitted in pediatric intensive care unit (PICU) with purpura fulminans.

**Patients and methods** Retrospective review of case sheets was done. Sixty-nine children (median age 3 years (1.6 months to 11 years and boy to girl ratio 1.5) admitted between January 2000 and May 2015 with PF in PICU of the pediatric hospital Bechar Hamza in Tunis, were included. The diagnosis of PF was made in patients with severe sepsis or septic shock with an extensive purpura. The PF is considered secondary to meningococcal infection if Neisseria meningitidis or soluble antigens are found at the blood culture or cerebrospinal fluid. In patients whose samples are negative, the PF is considered secondary to meningococcal infection if no other bacterial or viral origin is found to explain the purpura. For each patient enrolled, we have clarified the demographic data, the severity of clinical presentation by pediatric risk score of mortality (PRISM) and Glasgow meningococcal septicemia prognostic score (GMSPS) and therapeutic data.

**Results** A mortality rate of 52% was observed for a predicted death by the PRISM score at 24.6%. Seventy-seven percent of the deaths occurred during the first 24 h. Twenty-five (69.4%) children died of irreversible septic shock and 6 (16.6%) children died of refractory hypoxemia. Independent predictors of mortality were the initial severity assessed by the GMSPS (p = 0.001) and use of high doses of vasoactive drugs evaluated by the vasoactive-inotropic score (p = 0.026).

**Conclusion** The mortality in our PICU is high, higher than predicted by PRISM. It occurred mainly within 24 h of admission. Early recognition and prompt initial antibiotic therapy continue to be the cornerstones of the successful management of this dramatic disease, reducing mortality.

**Competing interests** None.
P70
Epidemiology multiresistant bacteria in pediatric medicosurgical intensive care unit
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Introduction Nosocomial infections (NI) to multi-resistant bacteria (BMR) is a main public health problem worldwide. They are particularly frequent and severe in pediatric intensive care.

Objectives To evaluate the incidence, bacteriological profile, epidemiological characteristics and resistance associated to BMR antibiotics acquired in pediatric intensive care unit.

Materials and methods Retrospective study including inpatients and spending more than 48 h in medical-surgical pediatric intensive care unit, at Ibn Rochd University Hospital of Casablanca, over a period of 12 months from 1 January 2015 to 31 December 2015. The BMR taken from different samples (PBDP, central catheter, hemoculture, LCR, urine culture and sampling devices).

Results During the study period, we collected 30 episodes of NI to BMR, the incidence rate was 7.1% and the incidence density was 20.6% per 1000 hospitalization days. Two infectious sites were preponderant: pneumonia (53.3%), and central catheter infections (40%). Gram-negative bacilli (GNB) resistant to C3G were most common (about 120 samples: 87%). Acinetobacter baumanii was the main species (60 samples: 43.5%). These GNB resistant to C3G had presented resistance exceeding 50% for imipenem, amikacin, gentamicin, tobramycin, and cotrimoxazole. Resistance to ciprofloxacin was 16.6%. They had isolated 30 strains of Pseudomonas aeruginosa and 24 klebsiella pneumonieae strains resistant to C3G. Six strains of Staphylococcus coagulase negative (SCN) were isolated 2 of which resistant to vancomycin. No strain was resistant to glycopeptides.

Conclusion BMR Nosocomial infections are frequent and serious in pediatric intensive care unit. Thus, continuous monitoring of resistance in nosocomial bacteria to antibiotics is a main concern.

Competing interests None.

P71
An outbreak of Serratia marcescens in a mixed neonatal and paediatric intensive care unit: investigation of causes and management
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Introduction S. marcescens, a gram negative bacillus, classified as an Enterobacteriaceae, was originally considered to be a non-pathogenic saprophytic water organism. Since several years, it is a well-recognised colonisation. All infected new-borns have favourable clinical outcome.

Patients and methods Since several years, it is a well-recognised saprophytic water organism. Since several years, it is a well-recognised colonisation. All infected new-borns have favourable clinical outcome.

Results Between October 2015 and February 2016, seven preterm new-borns (gestational age: 25–34 weeks) were identified with S. marcescens new-borns (gestational age: 25–34 weeks) were identified with S. marcescens.

Conclusion S. marcescens can cause rapidly-spreading outbreaks of severe infections in neonatal units, but with appropriate control measures these outbreaks can be contained at an early stage.

Competing interests None.

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evolution, were recorded. A paediatric infectious disease specialist reviewed the charts to determine for each patient if the modification or the remaining of initial treatment were relevant or not. An alternative choice was suggested in case of irrelevance.

Results We included 168 antimicrobial treatments. 87.4% were adjusted within 72 h after initiation. 77.3% of adjustments were considered relevant. Among those that had not been adjusted, de-escalation would have been possible in 92.9% of cases. Among all those which should have needed an adjustment, 26.1% had an insufficient or non-realised de-escalation. Factors associated with absence of or irrelevant adaptation were nosocomial infection and absence of consultation of a paediatric infectious disease specialist about antibiotic modification.

Conclusion Antimicrobial treatment adjustments are globally well executed in our paediatric intensive care unit, but could be improved by obtaining expertise from an infectious diseases specialist more frequently. Hence antibiotic de-escalation should be improved. The results of this study constitute an adequate basis for a future antimicrobial stewardship that could be conducted in our unit while focusing on a systematic consultation with a paediatric infectious disease specialist.

Competing interests None.

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P74 Does apnea affects the reliability of Analgesia Nociception Index (ANI), as a monitor of nociception?
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Introduction Continuous assessment of nociception intensity during anaesthesia is crucial. Analgesia nociception index (ANI: 0–100) has been proposed to clinicians to objectively measure pain in both conscious and anesthetized patient. ANI calculation results from sophisticated computing of sympathetic to parasympathetic tones balance using heart and respiration rate variability. Because we aimed to evaluate nociception during oro-tracheal intubation performed under anaesthesia conditions, we conducted a study to test the reliability of ANI monitoring nociception when respiratory rate was zero.

Materials and methods ASA physical status I healthy informed and consenting adult volunteers participated in this experimental prospective trial. Demographic parameters were recorded and non-invasive hemodynamics (HD), blood pressure (BP), heart rate (HR), pulsed oxygen saturation (SpO2) and ANI monitoring tools were installed. After a 5 min resting and stabilization the volunteers lying on their back were asked to perform a 30 s duration apnea starting on command at Residual Functional Capacity (RCF). After a wash-out period following this first apnea period (P1) and return to baseline parameters, the volunteers were asked again to perform a second 30 s duration apnea (P2) starting on command at RCF. Initiation of P2 coincided with 30 s unlar nerve stimulation at the wrist (single twitch, 1 Hz, 25 mA). The different component of ANI (energy, mean, and instantaneous values: iANI), HR, BP and SpO2 values were recorded at predefined time points: T0 (baseline values, just before apnea), T1 (end of apnea), T2 (T1 + 30 s), for P1, and the same predefined time points (T0', T1', T2') for P2. After the experiment, volunteers were asked to rate the pain intensity at the wrist during P2, using a Visual Analogue Scale (VAS: 0–100). Evolutions of measured parameters were compared between P1 and P2. Values are mean ± SD.

Results 21 healthy volunteers aged 34 ± 10 were included. During P1 and P2, both iANI and energy were monitored. Mean VAS during P2 was 27±100. Figure 23 illustrates HD and iANI variations at T1/T1' and T2/T2'. During P1, there is a remarkable stability of iANI at T1 (+1%) and no significant decrease at T2 (~3%). On the opposite, a progressive decrease of iANI was evidenced during P2 (~5% at T1'), reaching a nadir 30 s after the end of nociceptive stimulation (~17% at T2'). HD and SpO2 remained stable and comparable in P1 and P2.

Conclusion Based upon our results, ANI seems discriminant and efficient at measuring nociception in the conditions of a short apnea of 30 s. ANI is thus probably a reliable tool to measure nociception during oro-tracheal intubation. Our trial shows that HD is less sensitive than ANI to detect weak intensity nociceptive stimulus.

Competing interests None.

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P75 Early mobilization in a medical ICU: a first Tunisian experience
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Introduction Critically ill patients are known to be complicated and their mobilization on the earliest days of ICU stay has shown to be
feasible, well tolerated and beneficial. We aimed to assess the feasibility and tolerance of early mobilization as part of routine care in a Tunisian medical ICU.

**Patients and methods** A prospective period study conducted in a 7-bed-medical ICU with nurse/patient ratio at 1/3 over 12 months. We were studied, incidence density of four predefined activity events (sit on bed, sit in chair, stand-up and ambulate), onset delay, limitation factors and tolerance. Six activity-related adverse events were defined as fall to knees, catheters or tubes removal, systolic pressure drops more than 30 mmHg, pulse increase by 20 beats/min or bradycardia, SpO2 drop <80% and extubation.

**Results** During the study period, we conducted a total of 1265 activity events in 127 patients. The mean length of stay was 11.8 ± 12.3 days and the global incidence density was 842 activities/1000 patient days. The mean age was 56.4 years ± 19.9, 44(34) patients had a BMI > 30 kg/m². Mean SAPS II score was 29.9 ± 12.9 years. At ICU admission 54(42.5) patients were on invasive mechanical ventilation, 38(29.9) patients were on NIV and 27(21.3) patients had vasopressors. Onset delay to the first activity was 5.8 ± 3.8 days for patients on mechanical ventilation. 21(16.5) patients had vasopressors when they did their first activity and the onset delay for those patients was 5.6 ± 3.5 days. The first activities consisted only in “sit on bed”, 80.3% and “sit in chair”, 19.7%.

Neither standing nor ambulation was done as a first activity. The activity events included 306(24.2) sit on bed, 629(49.8) sit in chair, 182(14.3) standing position and 148(11.7%) ambulation. Incidence density of activities was 604 activities/1000 patient days for patients with invasive mechanical ventilation, 1233 for those with NIV and 1004 for patients without mechanical ventilation. In patients on ambulation 6 had endotracheal tube, 19 had tracheostomy and 27 had NIV. Twenty-eight incidents occurred (2.1%). They consist in hypotension (32.1%), tachycardia (46.4%), one fall to knees, 3 feed tube removals, one urinary tube removal and only one central catheter removal was registered without harmful consequences. No accidental extubation occurred. No barrier to critically ill patient was found.

**Discussion** Our study proves that early mobilization is feasible and safe for all patients including those with mechanical ventilation. Our results are in line with literature [1]. Sricharoenchai in an observational study conducted in Baltimore ICU over 30-month period including S267 activities showed a low incidence of early mobilization adverse effects (0.6%) [2].

**Conclusion** This study shows the feasibility and safety of an uncommon practice in a relatively poor resources and low patient-nurse ratio medical ICU.

**Competing interests** None.

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Table 17 Time data

| Time data                  | Team Config A | Team Config B |
|----------------------------|---------------|---------------|
| Number of interventions    | 2342          | 4287          |
| Percentage of interventions| 35.3          | 64.7          |
| Total duration of all interventions (h) | 1088 | 2575 |
| Percentage of duration of all interventions | 29.7 | 70.3 |
| Average time of intervention (min) | 28 | 36 |
| Maximal duration (min)     | 285           | 390           |

having consulted with medical emergencies over a week. The data collected were epidemiological and socio-economic, the reason for the consultation and the future of consultants. The consultations were deemed appropriate or not by the doctor on duty depending on the nature of the disease, day and time of the consultation and the age of the symptoms.

Results We collected 469 consultants. The average age was 43.45 years (16–104 years) for a sex ratio of 0.87. The main reasons were an opharyngeal infection or bronchopulmonary (17.3%), asthenia (13.4%), chest pain (11.9%). More than 50% of the consultations were deemed inappropriate. These patients were younger than 35 years in 48.4% of cases, and jobless in 58.6% of cases. There was no significant difference in the number of inappropriate consultations between the two genders. The days and times of arrival did not differ between appropriate consultation and not appropriate. After review, 6% of patients were admitted to unit short hospitalization, 49.9% referred to a specialist consultation, other consultants put outgoing, equipped (39.7%) or not (4.5%) of a therapeutic prescription.

Discussion The use of emergency would be linked to the absence of prior appointment, and continuity of care. The consultants have the option of availability, some for conditions not covered by the emergency. The high proportion of younger people without profession is explained by the free care. The elderly rarely autonomous and complex diseases carriers, are fewer in the “inadequate consultation” group.

Conclusion Develop a care system and quality of care in basic health facilities, educate and convince patients to get there, would relieve hospital emergency services that will then take care of true emergencies.

Competing interests None.

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P79 Impact of the evolution of the National Health Insurance tariffs on the income of a French ICU

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Introduction Hospital funding for ICU stays in France is made of reimbursement of a fixed amount according the diagnosis-related group (DRG) of the patients and of extra funding for each day spent in ICU if the patient fulfilled criteria of severity (SAPS2 > 15) and of treatment intensity (use of organ support). The tariffs of reimbursement for the different DRGs and for the extra funding are updated every year. We measured the impact of these updates on the income of our ICU.

Patients and methods DRG and length of stay of all the patients hospitalized during 2011 in our 12-bed ICU were extracted from the administrative database of our institution. We computed the reimbursement for the care of these patients with the tariffs of 2011, 2012, 2013, 2014, 2015 and 2016 (data from the Agence Technique pour l’Information Hospitalière). Inflation was taken into account according to the recommendations of the Institut national de la statistique et des études économiques.

Results 592 ICU stays (3224 days) have been analyzed. The patients were classified in 237 DRG. The income of the ICU decreased from 8,416,260, 14€ in 2011 to 7,816,786, 72€ (−7%). Income decreased every year, with the most important lessenings during the first years (2012: −2.37%; 2013: −2.47%; 2014: −0.81%; 2015: −0.82%; 2016: −0.84%). This reduction was explained by both a lowering of the tariffs of the different DRG (mean evolution −4.6%) and a diminution of the extra funding (814.32 € per day in 2011, 801.19 € in 2016, −1.6%).

Discussion These results are based on a small number of ICU stays but are probably representative of the global trends of ICU reimbursement because of the high number of DRG analyzed.

Conclusion This simulation gives an estimate of the economical pressure sustained by the French ICUs during the last 6 years. Productivity gains are necessary to cope the tariff evolution and may require reduction of costs, increase of activity and pricing optimization.

Competing interests None.

P80 Risk factors and prognosis of C sections performed in emergency for placenta previa bleeding

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Annals of Intensive Care 2017, 7(Suppl 1):P80

Introduction Hemorrhagic complications during placenta previa are unpredictable and may indicate an extremely urgent C section. Our study goal is to determine the risk factors and prognosis of this urgent situation.

Patients and methods This is a case–control study, prospective and analytic over the second semester of 2015. We included all patients who underwent emergency C section for placenta previa bleeding. The witness cases are represented by C section performed in emergency for another reason, at random.

Results We identified 42 cases for 126 witnesses. The over 30 years patients accounted for 45.23% in the case group against 22.2% in the controls (p = 0.03). The average rate for cases was 3–2 against the controls (p = 0.001). For instance, the admission was referred in 76.19% in the case group against 80.95% for controls. At least an ultrasound was performed during pregnancy in 23.8% of case group against 32.5% for control. Maternal complications in the case group are represented by a maternal anemia in 85.71% against 15.07% for controls (p = 0.002), fetal complications due to hypotrophy represent 19.04% in the case group against 11.1% for controls (p = 0.04), the perinatal mortality was 26.19% in the case group against 9.5% for controls (p = 0.03).

Conclusion The placenta previa is still a serious disease of pregnancy. Early ultrasound diagnosis would enable appropriate monitoring and prevention of all obstetric complications of this disease.

Competing interests None.
Unusual vein thrombosis
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Introduction Unusual venous thrombosis are those whose site is other than the lower limbs. They are quite rare. Clinical expressions depend on their locations (digestive, brain or upper limbs).
Their diagnosis and treatment are much less codified than the lower limbs locations. Their causes should be carefully sought for better management.

Materials and methods Retrospective and descriptive study of patient records who experienced deep vein thrombosis and were followed in the service during the period between 1985 and 2015. Our work aims to classify these atypical localizations in our patients, and to study the epidemiology, etiology, treatment, and finally their evolution.

Results We collected 120 cases of deep vein thrombosis, of which 25 cases are unusual location, it is 6 inferior vena cava thrombosis, 1 superior vena cava thrombosis, 8 thrombosis of renal vein, one thrombosis of hepatic vein, 4 cerebral localizations, 4 locations at the Superior Members, 1 thrombosis of the internal jugular vein and 2 of the subclavian vein and 2 of the axillary veins. One thrombosis of the central vein of the retina and one of thrombosis renal graft vein. There were 17 women and 8 men whose average age was 43.4 years.
The diagnostic tool was Doppler ultrasound in 60% of cases, CT or magnetic resonance imaging in 40% of cases.
The etiologic survey found that 7 patients had glomerular nephropathy with an acute nephrotic syndrome, 1 patient had lupus, 4 patients with Behcet's disease, 4 patients had neoplasia, 1 patient had protein C and S deficiency, 1 patient had hyperhomocysteinemia and no etiology was found in 1 case.
All patients were put on anticoagulants associated with etiological treatment. The outcome was favorable in 13 patients, 2 patients had pulmonary embolism, 2 patients died, in 3 patients thrombosis recurred and 5 patients were lost to view.

Conclusion In the contrary to the literature, the neoplastic origin of unusual thrombosis does not prevail in our series where the causes are dominated vasculitis and renal disease which is explained by the selection bias of patients.

Competing interests None.

Reference
1. Rev Internal Med.

P82

Pain therapy in the emergency: satisfaction survey
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Introduction The management of pain is an emergency medicine’s priority. It is a criterion of quality care. We conducted a survey that assesses the degree of patient satisfaction overlooked the pain therapy.

Patients and methods Prospective observational study performed in the emergency department, which included all patients aged over 10 years consulting for acute pain requiring the use of an analgesic treatment. The intensity of pain was measured by visual analogue scale (VAS). Patients unable to assess pain by VAS or having a vital distress are excluded from this study.

Results Over a period of 3 months, we included 183 patients. The average age was 41 ± 16 years. The sex ratio was 1.7. The mechanism of pain was traumatic in 25 patients (13.7%). VAS means at the entry was 5.8 ± 2.6. At the exit, it was 6.4 ± 2.3. We used the painkillers levels 1 in 145 patients (79.2%), levels 2 in 27 patients (14.8%) and levels 3 in 11 patients (6%). Only 6% of patients are not satisfied, 38.8% are dissatisfied, while 44.8% are satisfied or very satisfied (10.4%).

Conclusion This study shows that nearly 50% of patients are few or dissatisfied with the management of pain, efforts are still needed to improve.

Competing interests None.

Reference
1. Rev Internal Med.

P83

Transcranial Doppler in pediatric intensive care units for 152 children without traumatic brain injury
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Annals of Intensive Care 2017, 7(Suppl 1):P83

Introduction Transcranial Doppler (TCD) is currently used at the bedside of critically ill children and can detect cerebral blood flow modifications. Use and impact of TCD in non-traumatic critically ill children is unknown. We aimed to describe and to assess the TCD and related impact in this population.

Patients and methods This French prospective and multicentric study included all children (0–18 years) who had TCD in four pediatric intensive care units during 1 year. A questionnaire was completed for each performed TCD by the intensivist. TCD interpretation and impact were accorded to the intensivist. Factors associated with impact, and especially diagnostic and therapeutic impacts were identified.

Results 152 patients were included with a median age of 7.6 months (0–206 months). TCD was performed in 6.7% of all admitted patients during the inclusion period. 61% of patients were aged between 0–206 months. TCD was performed in 6.7% of all admitted patients during the inclusion period. 61% of patients were aged between 0–206 months. TCD was performed in 6.7% of all admitted patients during the inclusion period. 61% of patients were aged between 0–206 months. TCD was performed in 6.7% of all admitted patients during the inclusion period. 61% of patients were aged between 0–206 months. TCD was performed in 6.7% of all admitted patients during the inclusion period. 61% of patients were aged between 0–206 months. TCD was performed in 6.7% of all admitted patients during the inclusion period. 61% of patients were aged between 0–206 months. TCD was performed in 6.7% of all admitted patients during the inclusion period. 61% of patients were aged between 0–206 months. TCD was performed in 6.7% of all admitted patients during the inclusion period. 61% of patients were aged between 0–206 months. TCD was performed in 6.7% of all admitted patients during the inclusion period. 61% of patients were aged between 0–206 months. TCD was performed in 6.7% of all admitted patients during the inclusion period. 61% of patients were aged between 0–206 months. TCD was performed in 6.7% of all admitted patients during the inclusion period. 61% of patients were aged between 0–206 months. TCD was performed in 6.7% of all admitted patients during the inclusion period. 61% of patients were aged between 0–206 months. TCD was performed in 6.7% of all admitted patients during the inclusion period. 61% of patients were aged between 0–206 months. TCD was performed in 6.7% of all admitted patients during the inclusion period. 61% of patients were aged between 0–206 months.

Discrepancy between the interpretation of DTC values and published reference values was found in 116 patients (76%). TCD had an impact for 116 patients (76%), with a strong impact (diagnosis’ confirmation or exclusion, therapeutic impact) for 98 patients (64%). Confirmation of diagnosis was statistically associated with patients severity: mortality, catecholamines, intubation, PELOD score (p < 0.05). Hemodynamic dysfunction (OR 6.4; p = 0.007) and less experienced operators (OR 3.9; p = 0.012) were independent factors affecting therapeutic impact in multivariate analysis.

Discussion This original study is at our best of knowledge the first to describe use of TCD in heterogeneous diseases in critically ill children without traumatic brain injury. TCD was performed not only in patients with neurological disorders, highlighting the wide field of applications of this exam. This study showed the difficulty of TCD interpretation, whose reference values are not well-known by operators. Moreover, TCD values could be biased by several systemic parameters, usually not controlled in these unstable patients. TCD’s impact was
highly retained, and depending on severity of patients and on intensivist’s experience. 

**Conclusion** TCD is performed in heterogeneous diseases in non-traumatic critically ill children. TCD has an important impact, dependent on the operator’s experience and on patient’s severity. Regarding the difficulty of accurate interpretation of TCD’s values, improvement of TCD’s use is necessary.

**Competing interests** None.

**P84**

**Recruitment of the sublingual microcirculation in children with extracorporeal membrane oxygenation**

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**Annals of Intensive Care 2017, 7(Suppl 1):P84**

**Introduction** The sublingual microcirculation is impaired in states of severe sepsis and septic shocks. The microcirculatory dysfunctions are prognostic markers of survival and organ failures in critically ill adult patients. In children, the sublingual microcirculation is also impaired in septic shock [1] and severe respiratory failures [2]. We hypothesized that the microcirculation impairments related to the severity of respiratory and circulatory failures could be reversible under extracorporeal membrane oxygenation (ECMO). The aims of the study are (1) assess the feasibility of the sublingual microcirculation monitoring in pediatric ECMO, (2) compare the microcirculation parameters under ECMO between J1 and J3.

**Patients and methods** Prospective observational study in the pediatric intensive care unit at Trousseau hospital, Paris. The videos were acquired with Microscan® device (Microvision, Amsterdam) in sublingual areas. The data analyses have been performed by the AVA3 software. The statistical analyses have been performed by PRISM 5 software.

**Results** We have included 10 children on ECMO. The age and weight were respectively 5 ± 14.8 months [0;47] and 4.7 ± 4 kg [2.5;16]. The sex ratio (M/F) was 3/7. The ECMO indications were refractory acute respiratory distress syndromes (4), septic shocks (3), cardiac arrest (1), diaphragmatic hernias (3), refractory pulmonary hypertension (1). During the pre-ECMO period 9 children have been treated with Nitric Oxide for pulmonary arterial hypertension and 9 children needed catecholamine support. During ECMO 8 children have been treated with noradrenaline, 3 with adrenaline, 6 with dobutamine. The mean catecholamines duration was 6.5 ± 8.2 days [0;25]. Five children received a arterio-venous ECMO, 2 veno-venous ECMO, and 3 multimodal ECMO (A-V and V-V). The mean ECMO duration was 9.4 ± 3.3 days [2;13] and hospitalization duration was 29.6 ± 20 days [9;64]. Concerning the microcirculation parameters the microvascular flow index (MFI) was significantly improved in the small microvessels (2.1 ± 0.36 [1.3; 2.5]) at day 1 and 2.5 ± 0.27 [2.2; 3] at day 3; p = 0.00173) and in the medium microvessels (2.6 ± 0.24 [2.2; 2.9] at day 1 and 2.8 ± 0.2 [2.5; 3] at day 3; p = 0.045). Regarding the biological markers the pH increased between day 1 and day 3 (7.3 ± 0.12 [7.1, 7.5] vs 7.4 ± 0.11 [7.3, 7, 6]; p = 0.00645. See Fig. 24.

**Discussion** In this preliminary study the children had severe hemodynamic failures or severe respiratory failures with pulmonary hypertension which needed of catecholamines and ECMO support. The alteration of the sublingual microcirculation was confirmed at day 1 under ECMO. The microvascular perfusion parameters were reduced compared to the normal value of healthy subjects (MFI = 3). The partial MFI restoration confirmed the microcirculation recruitment in these patients. Now, further studies are needed to confirm this microcirculation’s recruitment in a larger pediatric ECMO cohort and to investigate the impact of this recruitment on the prognosis.

**Conclusion** The microcirculation was impaired in the initial phase of ECMO in children with severe respiratory or circulatory failures and it was partially restored during the early phase of ECMO.

**Competing interests** None.

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**P85**

**Impedance cardiography by PhysioFlow® for non-invasive cardiac output monitoring: a comparison with trans-thoracic echocardiography in pediatric intensive care patients**

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**Annals of Intensive Care 2017, 7(Suppl 1):P85**

**Introduction** Impedance cardiography (IC) is a promising non-invasive, continuous cardiac output (CO) monitoring method. PhysioFlow® (PF®) is a new IC technique never studied in pediatric intensive care unit (PICU). The aim of the study was to compare CO and cardiac index (CI) measurements in PICU patients by IC using PF® with those obtained by the method of reference, trans-thoracic Doppler echocardiography (TTE), with regard to accuracy, precision of agreement and reproducibility.

**Patients and methods** In this single-center prospective method comparison study, all PICU patients aged between 28 days and 10-year-old and requiring TTE were included, except those with complex congenital heart disease or poor IC signal and/or TTE quality. Simultaneous sets of three measurements were realized by TTE and PF®. CO and CI measured by TTE (COTTE and CITTE) were compared with CO and CI obtained by PF® (COIC and CIC). Concordance correlation coefficient (CCC) and Bland–Altman analysis were used to analyze the concordance rate and compare accuracy and percentage error (PE) between the two methods. As required, data were logarithmically transformed prior to Bland–Altman analysis. Reproducibility was evaluated with intraclass correlation coefficient (ICC) and using the calculation of the precision of the method [1]. Post-hoc signal analysis was performed to evaluate the quality of IC signal.

**Results** A total of 43 patients (median age: 13 months, interquartile range (IQR) 4–34 months) were included. Median PIIM2 probability of death was 1.04 (IQR 0.34–5.11). 9.3% of patients had inotropic support and 30.2% had a mechanic ventilation. On 129 paired measurements, mean COTTE was 1.74 ± 0.93 L/min whereas mean COIC was 2.23 ± 1.21 L/min. Concordance for CO measurements was considered

![Fig. 24 Time course of the sublingual microcirculation and pH during ECMO. MFI The microvascular flow index evaluates the microvascular perfusion. *p < 0.05, **p < 0.005](image-url)
as good, since CCC was $r = 0.54$ (95% CI: 0.43–0.63). However, mean absolute bias for CO was 0.80 L/min (40%) with an unacceptable PE of 158%. Concerning CI measurements, mean CITE was 3.74 ± 1.00 L/min/m² whereas mean CIIC was 4.93 ± 1.95 L/min/m². Concordance for CI measurements was low, since CCC was $r = 0.12$ (95% CI: −0.08 to 0.31). Mean absolute bias for CI was 0.77 L/min/m² (18%) with an unacceptable PE of 62%. Nevertheless, reproducibility of PF® for CO and CI measurements was very good: ICC were $r = 0.94$ and $r = 0.90$ for CO and CI, respectively. Precision of PF® for CO measurements was 6.9% (IQR 4.6–11.6%) and 7.3% for CI (IQR 4.2%-12.2%). Age, sex, weight, heart rate, and hematocrit didn’t affect differences of CO and CI between the two methods. Post-hoc signal analysis revealed that only 67.4% patients had an acceptable quality for IC signal, and PF®’s algorithm didn’t recognize adequately the whole IC signal in 37.2% of our pediatric patients.

**Conclusion** In this first method comparison study testing CO monitoring by IC with PF® in PICU, PF® can’t accurately estimate CO and CI in comparison with TTE, mainly because of signal analysis and algorithm failure. Nevertheless, PF® has a very good reproducibility for CO and CI measurements. Trending ability of PF® should be tested, and this device could be able to monitor fluid challenge response.

**Competing interests** None.

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Factors associated with extracorporeal membrane oxygenation treatment in congenital diaphragmatic hernia: 10 years of experience
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Annals of Intensive Care 2017, 7(Suppl 1):P88

Introduction Congenital diaphragmatic hernia is a rare anomaly that occurs in <1–5 of every 10,000 births. Pulmonary hypoplasia and pulmonary hypertension are commonly associated with congenital diaphragmatic hernia and explain its high morbidity and mortality. Despite significant advances in neonatal intensive care and protective ventilation, mortality remains as high as 50%. The development of extracorporeal membrane oxygenation (ECMO) support has improved the outcome but its use is controversial in this pathology. The main objective of this study is to determine associated factors with extracorporeal membrane oxygenation (ECMO) in congenital diaphragmatic hernia and to explore ECMO’s indications, causes of death, and evolution after discharge for survivors.

Patients and methods We led a retrospective study in all neonates born with congenital diaphragmatic hernia and admitted to our institution between 2003 and 2015. Antenatal data, including observed/expected foetal pulmonary volume by magnetic resonance imaging, data of the first day in paediatric intensive care unit before ECMO initiation, complications, causes of death, characteristics of ECMO group, surgical parameters and evolution of survivors after 1 month of life were recorded in the national database for congenital diaphragmatic hernia and patients’ charts.

Results Among the 62 cases of congenital diaphragmatic hernia, 50 infants were included with 14 (36%) requiring ECMO. Global survival rate at 1 month of life was 39/50 (78%). Lowest observed/expected lung foetal volume (p = 0.016), lowest Apgar at 1 min (p = 0.041), highest oxygenation index (p < 0.001), highest SNAP–PE 2 score (p < 0.001), left ventricle dysfunction (p = 0.046), highest lactatemia (p = 0.002), highest PaCO2 (p = 0.007) were significantly associated with ECMO support. Among 6 (17%) deaths in the non-ECMO group, 5 (83%) were due to absolute contraindications to ECMO. Complications occurred more frequently in the infants requiring extracorporeal membrane oxygenation support.

Conclusion Associated factors for the need for ECMO in congenital diaphragmatic hernia have been identified. They should be useful to determine earlier the need for ECMO and prognosis for counselling parents.

Competing interests None.

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Diagnostic accuracy of abdominal compression for predicting fluid responsiveness in children
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Introduction Administration of fluid to increase cardiac output is a cornerstone of the hemodynamic resuscitation. The respiratory variation in aortic blood flow peak velocity (ΔVpeak) is the only variable to reliably predict fluid responsiveness in children. It requires assist control ventilation with high tidal volume (>10 ml/kg) [1]. The purpose of this study was to evaluate the clinical usefulness of assessing variation of stroke volume during a calibrated abdominal compression for the prediction of fluid responsiveness in children during acute circulatory failure.

Patients and methods This study was approved by local institutional review boards (CPP Lyon Sud Est III). Written informed consent was obtained from parents. Oral consent of patients old enough to understand the study was also obtained. Patient, less than 8 years old of two paediatric intensive care unit (PICU), during a 1-year period inclusion, from September 2015 to September 2016 were selected. Stroke volume index was assessed with an echocardiography at baseline, after an abdominal compression (with a calibrated pressure of 25 mm Hg), at a return to baseline, and after a volume expansion (10 ml/kg of fluid challenge over 10 min). Pulse pressure (PP) Systolic (SP), diastolic (DP), and mean arterial pressure (MAP); heart rate (HR), respiratory aortic blood flow velocity (ΔVpeak), left ventricular ejection fraction (LVEF), and respiratory vena cava diameter variation (VCvar) were also recorded before volume expansion. Stroke volume index was calculated as the left ventricular outflow tract (LVOT) surface multiplied by the LVOT Velocity time integral (VTI). Patients were classified as responders to fluid loading if their stroke volume index (SVI) increased by at least 15%. R software with pROC package was used to perform descriptive and analytic statistic. Pearson correlations were performed and Receiver operative characteristic curves were built. Bootstrap technic was used to compute confidence interval (CI). p < 0.05 was considered significant.

Results Thirty-one children were included, 17 (±2) month old and weighing 8 (±5) kg. Seven were not on mechanical ventilation and 15 were in a mode allowing spontaneous breathing. 16 patients were fluid responders and 15 non-responders. All the echocardiography were performed by MJL and NT. Operators were blind from the value of the VTI for each condition. Coefficient of variation of the SVI was 8.4 (CI 5.4;11.4)% and the least significant change of five averaged SVI was 10.6 (CI 6.8–14.5)%). Changes in SVI during abdominal compression and after a fluid challenge were correlated (R2 = 0.796; p < 0.001). Change in SVI and respiratory vena cava diameter variation or delta Peak before abdominal compression were not significantly correlated with change.
in SVI during volume expansion. PP variation and MAP variation during abdominal compression were not correlated with SVI variation after fluid challenge. The ROC curve analysis showed that SVI change during abdominal compression predicts fluid responsiveness. (AUC ROC = 0.93; CI: 0.82–0.99). The best threshold was 9.28% with a sensitivity 75% (CI: 0.50–0.94) and a specificity of 93% (CI: 0.79–1.00). AUROC of ΔSVI was 0.70 (CI: 0.48–0.91). AUROC of ΔVpeak was 0.58 (CI: 0.34–0.81).

Discussion Fluid responsiveness assessment, in children especially, is challenging when spontaneous breathing is authorized. It represents the vast majority of children hospitalized in PICU for acute circulatory failure. Abdominal compression is an old technic used to modify preload. Historically assessed with arterial pressure, it has never been formally evaluated so far. Our method seems quite reliable to predict fluid responsiveness, it was shown to be superior to SVIvar and ΔVpeak in our settings.

Conclusion SVI variation during abdominal compression was the sole reliable method to predict fluid responsiveness in a mixed population of children, with and without spontaneous breathing, suffering from acute circulatory failure.

Competing interests None.

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P90
Epidemiology of pediatric and adult fatal anaphylaxis in France: analysis of the French national data
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Annals of Intensive Care 2017, 7(Suppl 1):P90

Introduction Data on fatal anaphylaxis in France are limited. The aim of this study was to define the anaphylaxis mortality rate (AMR) in France between 1979 and 2011, in general and pediatric populations, and to examine the repartition by age, sex, cause and geographical regions.

Patients and methods All deaths in France are recorded by a physician and death certificates are collected and analyzed by the CEPIDC Unit (National Institute of Health and Medical Research, INSERM). Each death certificate records primary and secondary causes of death, demographic data including sex, age, birthday and geographic region of death. Anaphylaxis related deaths were identified by using the International Classification of Diseases (ICD) codes on the death certificates.

Databases of the French National Institute for Economical and Statistical Studies (INSEE) are available freely (www.insee.fr). Data regarding the characteristics of the French population, by region, year, sex and age were collected.

AMR were expressed per million persons and per year. All regression analyses were adjusted for age-effect and a multivariable log-linear Poisson regression model was further performed by including all of the predictors.

Results There were 1603 anaphylaxis related deaths in metropolitan France between 1979 and 2011, with 39 deaths in pediatric population (age less than 20 years). AMR was lower in pediatric population (0.08 per million population per annum, 95% CI 0.05–0.1) than in general population (0.85 per million population per annum, 95% CI 0.79–0.88) (p < 0.01). In general population, AMR was higher in male sex (1.08 per million population per annum, 95% CI 1–1.6) than in female sex (0.86 per million population per annum, 95% CI 0.78–0.89) (p < 0.01). Annual percentage change for case fatality rate was —2% (95% CI —2.5 to —1.5) indicating a decrease during the study period (p < 10–4). Latrogenic cause was the most common (63%), followed by «unspecified» (23%), venom (14%) and alimentation (0.6%). AMR was the highest in persons aged >70 years (3.50 [95% CI 3.25–3.76]) and the lowest in children. Venom-induced mortality rate was higher in the South region (0.16) compared to the North (0.11) (p = 0.003). Only 8 food-induced fatalities were recorded (age < 32 years in 7). With the multivariate analysis, older age and male sex were associated with an increased risk for anaphylaxis death of any cause (p < 10–4).

Conclusion Fatal anaphylaxis decreased in France between 1979 and 2011. Higher rates of fatal anaphylaxis are observed in male sex, group aged 70 years and older and iatrogenic cause. In pediatric population, fatal anaphylaxis is low, but it’s probably under-estimated by diagnostic difficulties.

Competing interests None.

P91
The burn out in the anesthesiology department
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Annals of Intensive Care 2017, 7(Suppl 1):P91

Introduction The burn out is an emotional state of exhaustion and loss of performance in response to chronic work stress. This study aims to assess the prevalence and determine the Burn out of risk factors in medical and paramedical staff in the anesthesiology department.

Patients and methods This was a multicenter, transversal analytic study performed within the medical and paramedical staff of the operation rooms in the central anesthesiology department in IBN ROCHD hospital in CASABLANCA. Caregivers were asked to freely and anonymously complete a questionnaire involving demographic variables, professional and an assessment of the causes and consequences of stress at work.

Results On a population composed of 60 caregivers, 41.7% had a score of high emotional exhaustion, 11.98% a high score of personalization and 47.38% a low score of professional achievement. Le burnout was found at 62.52% of our caregivers: 39.62% of them had low levels of burnout; 21.34% presented a moderate level and 1.58% had the highest level of burn out. In multivariate analysis, Doctors and Nurses Residents were most at risk of burn out. Fear of malpractice and the unsatisfactory salary multiplied the risk of burnout, respectively 2.14 and 1.83.

Conclusion Burn out is a threatening reality of the anesthesiology environment. The consequences could be severe on the personal and professional performance of the health institution, which involves the implementation of preventive strategies highlighting the value of work organization and the contribution of the and paramedical medical staff.

Competing interests None.

P92
Family satisfaction: impact of moving in new buildings
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Annals of Intensive Care 2017, 7(Suppl 1):P92

Introduction Evaluation of family satisfaction has become an essential part of the responsibility of intensivists and a major criterion for assessing the quality of care. Family satisfaction could depend on
Introduction The improvement of patient safety and quality of health care is the main goal of the Joint Commission International standards. One of these sections covers medication management and use. The standard recommends the requirement for a trained professional to review medication orders or prescriptions for appropriateness. In practice, this review process is usually performed by a clinical pharmacist. The aim of our study was to evaluate the impact of the clinical pharmacist intervention in medication orders.

Patients and methods This prospective study was conducted in a 7-bed medical ICU in Sousse, between November 2014 and August 2015. Patients who were admitted for at least one overnight stay were included. This study was performed in two periods, 1/an audit to identify the discrepancies of the medical prescriptions, 2/a “before/after” intervention study to evaluate the impact on medical prescriptions. The French Society of Clinical Pharmacy sheet was used to collect data.

Results Over a 10 months’ period, 68 patients were followed and 379 sheets were analyzed, gathering all the pharmacological classes. 148(39) discrepancies were identified, mainly related to drug interactions (69 (46.6)). 42% had no clinical impact, 42% had mild and none had severe impact. The most common classes involved in medication discrepancies were anti-infective, 53 (35.8); cardiovascular, 43 (29); systemic hormonal preparations, 14 (9.5) and anticoagulants, 13 (8.7). The post intervention period was addressed to the anti-infective drugs' prescription. The discrepancies rate was significantly reduced from 53 (63) to 33 (39) respectively in a total of 84 patients.

Conclusion Discrepancies seem to be common in a medical ICU, involving mainly drug interactions with rather a mild clinical impact. This could be significantly reduced by the intervention of a clinical pharmacist.

Competing interests None.

Impact of human factors on improving the management of the syndrome hellp
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Fig. 26 Comparison of CCFNI scores between 2009 and 2016
Table 18 Features of management

|                | Population 1 (N = 92 patients) | Population 2 (N = 112 patients) |
|----------------|--------------------------------|----------------------------------|
| AWT            | 30 min                         | 15 min                           |
| ATM            | 49 min                         | 24.5 min                         |
| MTBC           | 4 h 30                         | 2 h 30                           |
| AHS            | 29 days                        | 20 days                          |

AWT average waiting time, ATM average time management, MTBC mean time to blood components, AHS average hospital stay.

The average waiting time (AWT): Average time between the arrival of the patient and her admission.
The average time management (ATM): Average time between hospitalization and administration of the first treatment.
The mean time to blood components (MTBC): Average time between the application and administration of blood products.
The average hospital stay (AHS).

Results Records of these women were studied. The characteristics of women are the same for both populations, but there is a net reduction of maternal mortality 0–4 and the satisfaction rate increased from 31 to 71%. The features of management are summarized (see Table 18).

Discussion Our institution demand for unscheduled care and frequency of emergencies are increasing in the service of emergency obstetric gynecological similar phenomenon reported in some publications. Our study marks a reduced waiting time in half, as in a study published in France [2]. Despite the protocol established for the second population, we note a major impact for the average time of care, and the average time to obtain blood components is as long, related organizational deficits at night and on weekends, but this has allowed us to reduce the mortality rate and duration so stay on the economic cost.

Conclusion The human factor has a direct impact on the length of hospitalization and thus the economic cost, on maternal mortality, an indirect impact on satisfaction patient. It is time to realize that the human factor is the success factor.

Competing interests None.

Table 19 Evaluation of pre-formative knowledge: rate score above average

|                | T1 | T2 | T3 | T4 | T5 | T6 | T7 | T8 | T9 | T10 | T11 | T12 | T13 | T14 | T15 | T16 | M  |
|----------------|----|----|----|----|----|----|----|----|----|-----|-----|-----|-----|-----|-----|----|
| Nurse (%)      | 11 | 0  | 60 | 20 | 10 | 20 | 10 | 50 | 54 | 83  | 27  | 75  | 90  | 90  | 33  | 91  | 45 |
| Technicien (%) | 0  | 75 | 75 | 100| 0  | 25 | 100| 75 | 100| 100 | 33  | 100 | 100 | 66  | 66  | 66  | 67 |
| Healthcare assistant (%) | 0 | 25 | 50 | 0  | 0  | 25 | 25 | 33 | 33 | 0   | 75  | 100 | 50  | 50  | 100 | 33 |

P96 Involvement of paramedical staff newly recruited in intensive care in their own training

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Annals of Intensive Care 2017, 7(Suppl 1):P96

Introduction In the absence of specific academic training in intensive care for paramedical personnel and in order to improve skills, training is necessary especially before the opening of a new unit. The aim of our study was to evaluate the training of newly recruited in intensive care paramedical staff.

Patients and methods For initiation of a new intensive care unit, medical staff prepared a specific training program of resuscitation. The schedule of the training has been distributed to all relevant staff in advance.

Presentations (basics and technical data sheets) on various resuscitation themes were prepared by the staff and reviewed by doctors’ leaders. The evaluation form contained 16 tests (T). Each test consisted of five questions before the presentation and five after.

An increase in the evaluation mark obtained after the presentation is considered an improvement in knowledge.

Results Twenty of the staff: twelve nurses, four technicians in intensive care and four healthcare assistant participated in the training. Fifty-five percent of the staff were working in the private sector before their recruitment. The median time between graduation and recruitment was 1 year and a half. Technicians group had the highest level of knowledge (Table 19).

The analysis of the post-formative evaluation noted an increase in knowledge. The healthcare assistant group had the lowest rate of improvement (Table 20).

Conclusion The direct involvement of paramedical staff in their own training increases their motivation without harming the quality and the main objective: knowledge improvement.

Competing interests None.

Table 20 Knowledge improvement percentage

|                | T1 | T2 | T3 | T4 | T5 | T6 | T7 | T8 | T9 | T10 | T11 | T12 | T13 | T14 | T15 | T16 | M  |
|----------------|----|----|----|----|----|----|----|----|----|-----|-----|-----|-----|-----|-----|----|
| Nurse (%)      | 60 | 55 | 70 | 80 | 50 | 60 | 90 | 75 | 70 | 36  | 66  | 63  | 30  | 77  | 41  | 58  | 63 |
| Technicien (%) | 100| 25 | 75 | 25 | 25 | 75 | 50 | 25 | 66 | 66  | 66  | 100 | 100 | 33  | 33  | 58  |
| Healthcare assistant (%) | 75 | 50 | 75 | 66 | 0  | 100| 66 | 25 | 25 | 66  | 100 | 66  | 0   | 25  | 25  | 47 |
to assess the burnout prevalence in newly recruited paramedical intensive care staff before and after theory and practical training.  

**Patients and methods** Before the opening of a new intensive care unit, three questionnaires were filled by the paramedical staff: at the taking up, at the end of theory training and 3 months after practical training in intensive care units. The level of burnout was assessed using the “Maslach Burn Out Inventory” score and the degree of depression with Major Depression Inventory (MDI) test.  

**Results** Twenty of the paramedical staff: twelve nurses, four technicians in intensive care and four healthcare assistants participated in the study. Only one questionnaire has not been analyzed, it belonged to a nurse presenting a serious mood disorder. Forty-five percent of the study population was unemployed before their recruitment. Eight individuals (40%) showed burnout the first day of work, 11 (55%) at the end of training and 12 (60%) after 3 months of training (Table 21).  

Regarding the sub-dimensions of burnout there was an improvement of professional fulfillment (increasing from 35 to 45% after training and practicum). We also noticed a worsening of emotional distress (increasing from 0 to 25%). There was also an improvement in mood disorders (decreasing of 35–15%).  

**Conclusion** Theoretical and practical courses enhanced the level of professional achievement but were not sufficient alone to prevent Burnout.  

**Competing interests** None.

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**P98**

Does serum procalcitonin predict the onset of toxic acute hepatitis in acetaminophen poisoning?

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Introduction Procalcitonin (PCT) is a pro-hormone mainly produced by the thyroid C cells and routinely used as diagnostic biomarker of bacterial infection. PCT synthesis has never been described in the liver. To our best knowledge, no study investigated the possible predictive value of PCT in acetaminophen poisoning. Our objectives were to report the distribution of serum PCT values in acetaminophen-poisoned patients according to the onset and severity of their toxic liver injury in order to assess any possible predictive value for this biomarker.  

**Patients and methods** We conducted a retrospective single centre observational study including all acetaminophen-poisoned patients (either accidentally or voluntary) admitted to the ICU from 2013 to 2016. Patients were treated with the 3-bag N-acetylcysteine protocol according to the international recommendations based on the interpretation (when possible) of the plasma acetaminophen concentration on the Rumack-Matthew nomogram (line to treat the patient starting at 150 mg/L at the 4th hour). Serum PCT was measured using an automated method (Elecsys® and Cobase® analyzers; range: 0.02–100 ng/mL) and plasma acetaminophen concentrations were determined using spectrophotometry. Comparisons were performed using Chi-2 and Mann–Whitney tests.  

**Results** Seventy patients (50F/20M; age: 34 years [21; 53] [median [25; 75 percentiles]]; poly-intoxications: 83%) were included in the study. The presumed ingested acetaminophen dose was 15.5 g [8.0; 29.0]. The delay between acetaminophen ingestion and N-acetylcysteine infusion was 4.5 h [2.9; 9.0]. Serum PCT was markedly increased above the 1 μg/L threshold in the patients who already presented or further developed significant liver cytolysis defined by serum alanine ami- notransferase (ALAT) >100 UI/L (2 N) despite the treatment with N-acetylcysteine and independently of the onset of any bacterial infection, with a specificity of 97.9% and sensitivity of 69.6%.  

**Conclusion** Serum PCT measurement in acetaminophen-poisoned patients admitted to the ICU is helpful to early identify patients who present significant acetaminophen-related liver toxicity already established or in progress on admission despite the administration of N-acetylcysteine according to the international recommendations.  

**Competing interests** None.

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**P99**

Venlafaxine poisoning in the intensive care unit: clinical presentation and role of the cytochrome P450 2D6 phenotype in the onset of cardiovascular complications

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Introduction Venlafaxine, an antidepressant drug with properties of serotonin and noradrenaline reuptake inhibition, may be responsible in overdose for life-threatening cardiovascular complications. Venlafaxin-related toxicity additionally includes a possible interaction at elevated concentrations with the membrane sodium channels resulting in membrane stabilizing effects. To date, vulnerability factors to develop such cardiovascular complications are unknown. Based on a limited number of reported cases, poor cytochrome P450 (CYP) 2D6 metabolizers have been suggested to develop increased cardiovascular toxicity. This liver enzyme metabolizes venlafaxine to O-desmethyl-venlafaxine (or norvenlafaxine) and altered norvenla- faxine-to-venlafaxine metabolic ratio was suggested to support cardiac toxicity onset. We aimed to describe venlafaxine-related toxicity in patients admitted to the intensive care unit (ICU) and test the proposed hypothesis of vulnerability.  

**Patients and methods** We conducted a prospective single centre observational study including all venlafaxine-poisoned patients admitted to the ICU from 2010 to 2016. Plasma venlafaxine and norvenlafaxine concentrations were determined using gas chromatography coupled to diode array detector and mass spectrometer (LC-DAD/ MS). CYP2D6 genotyping was performed with the patient’s consent, allowing the description of venlafaxine-related toxicity in patients admitted to the intensive care unit (ICU) and test the proposed hypothesis of vulnerability.

**Results** Fifty-two patients (60% F/40% M; age: 44 years [32; 52], median [25; 75 percentiles]) exposed to venlafaxine (presumed ingested dose: 1.9 g [1.0; 3.0]; plasma venlafaxine concentration on admission: 0.8 mg/L [0.3; 2.0] and at the peak: 0.9 mg/L [0.4; 2.7]; 98% poly-intoxications) were included. Clinical features included consciousness impairment (Glasgow coma score: 8 [4; 14]) and seizure onset (14%), requiring mechanical ventilation (56%). Nineteen patients (37%) presented cardiovascular complications and three patients (6%) died in the ICU. Based on a univariate analysis, onset of cardiovascular toxicity was significantly associated with deeper coma (p = 0.04), reduced PaO2/FiO2 ratio (p = 0.004), onset of acute renal failure (p = 0.02), requirement of mechanical ventilation (p = 0.02)
and fatality (p = 0.04). No statistical relationships were found between cardiovascular toxicity and plasma venlafaxine concentration and norvenlafaxine-to-venlafaxine ratio on admission and at their respective peaks. When focusing on the patients with cardiovascular manifestations strictly attributed to venlafaxine toxicity, no statistical link was found with CYP2D6 phenotype.

**Conclusion** Venlafaxine poisoning may result in severe complications including cardiovascular toxicity and even fatality. Cardiac toxicity is responsible for increased morbidity-mortality but is not related to CYP2D6 phenotype. However, inclusion of additional patients is still warranted in our possibly underpowered study before any definitive conclusion.

**Competing interests**
None.

**P100**

Life threatening suicide attempts in Tunisia: an epidemiological and prognostic study

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**Annals of Intensive Care 2017, 7(Suppl 1):P100**

**Introduction** Suicide attempts represent a major public health problem in Tunisia according to recent data published in 2015 by the Tunisian National Social Observatory. However, data about the incidence of life threatening suicide attempts requiring intensive care unit (ICU) admission are lacking. The aim of this study is to describe the epidemiological profile of critically-ill patients admitted for suicide attempts and to identify factors predicting prolonged ICU stay (>3 days).

**Patients and methods** We conducted a retrospective study in the medical ICU of Bizerte city (Tunisia). From January 2009 to December 2015, 219 patients were consecutively admitted for suicide attempt. Three patients were excluded because of lacking data and 216 cases were included in the study.

**Results** The median age was 23 years [12–69]. The mean IGS2 score were 20.5 ± 14.5 and the mean APACHE2 was 8.1 ± 7. Psychotropic drugs overdose (43.1%) and pesticides ingestion (25.5%) were the two most common causes of suicide attempts. Eighty-one patients (37.5%) had a psychiatric history. Coma (32.9%) and respiratory distress (22.7%) were the two major reasons for ICU admission. Mechanical ventilation was required for 84 patients (38.9%). Median duration of mechanical ventilation was 2 days [1–9]. Overall mortality was 3.2%. The median ICU length of stay for the 209 survivors was 2j [1–19]. Factors associated with prolonged ICU stay (>3 days) were coma, hemodynamic instability, hypoxia, aspiration and mechanical ventilation.

**Conclusion** The current study is the largest study in Tunisia describing the epidemiological features of critically-ill patients admitted for suicide attempts. Identifying the underlying risk factors leading to suicide attempts and psychiatric follow-up starting upon ICU discharge are of paramount importance.

**Competing interests**
None.

**Reference**
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**P101**

Neurobehavioral effects of lithium in the rat: investigation of the effect/concentration relationships and the contribution of the poisoning pattern

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**Introduction** Severity of lithium poisoning depends on the ingested dose, previous treatment duration and renal function. No animal study has investigated neurobehavioral differences in relation to the lithium poisoning pattern observed in humans, while differences in lithium pharmacokinetics have been reported in lithium-pretreated rats mimicking chronic poisonings with enhanced brain accumulation in rats with renal failure. Our objectives were: (1) to investigate lithium-related effects in overdose on locomotor activity, anxiety-like behavior, spatial recognition memory and anhedonia in the rat; (2) to model the relationships between lithium-induced effects on locomotion and plasma, erythrocyte, cerebrospinal fluid and brain concentrations previously obtained according to the poisoning pattern.

**Materials and methods** Open-field, elevated plus-maze, Y-maze and sucrose consumption tests were used. We developed acute (intraperitoneal administration of 185 mg/kg Li2CO3 in naive rats), acute-on-chronic (intraperitoneal administration of 185 mg/kg Li2CO3 in rats receiving 800 mg/l Li2CO3 in water during 28 days), and chronic poisoning models (intraperitoneal administration of 74 mg/kg Li2CO3 during 5 days in rats with 15 mg/kg K2Cr2O7-induced renal failure).

**Results** In acutely lithium-poisoned rats, we observed horizontal (p < 0.001) and vertical hypolocomotion (p < 0.0001), increased anxiety-like behavior (p < 0.05) and impaired memory (p < 0.01) but no altered hedonic status. Horizontal (p < 0.01) and vertical (p < 0.001) hypolocomotion peaked more markedly 24 h after lithium injection and was more prolonged in acute-on-chronically versus acutely lithium-poisoned rats. Hypolocomotion in chronically lithium-poisoned rats with impaired renal function did not differ from acutely poisoned rats 24 h after the last injection. Interestingly, hypolocomotion/concentration relationships best fitted a sigmoidal Emax model in acute poisoning and a linear regression model linked to brain lithium in acute-on-chronic poisoning.

**Conclusion** Lithium overdose alters rat behaviour and consistently induces hypolocomotion which is more marked and prolonged in repeated lithium-treated rats. Our data suggest that differences between poisoning patterns regarding lithium-induced hypolocomotion are better explained by the duration of lithium exposure than by its brain accumulation.

**Competing interests**
None.
Mean duration of MV was 17 ± 4 h. The outcome was favourable in all cases. Length of hospital stay was less than 24 h with an average of 22 h [20, 24]. There is no correlation between coma and assumed ingested dose.

**Conclusion** Cyproheptadin overdose causes mainly neurological effects resulting from anticholinergic syndrome. Outcome is generally favourable.

**Competing interests** None.

### P103

**Cardiogenic shock complicating acute beta blockers intoxication: epidemiology and treatment**

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*Annals of Intensive Care 2017, 7(Suppl 1):P103*

**Introduction** In spite of its rarity, the acute Beta blockers (BB) intoxication represents a major cause of mortality as a result of cardiovascular toxicity. (CS) is the most serious complication of this intoxication. The objectives of this study were to determine epidemiological, clinical, management and prognosis of cardiogenic shock (CS) complicating acute BB intoxication.

**Patients and methods** This was a retrospective descriptive study conducted at the Medical Intensive Care Unit of CAMU over a 10 years period from January 2006 to December 2015. All hospitalized patients for cardiogenic shock complicating pure acute BB intoxication were included.

**Results** Twenty-five patients were included. The average age was 23 ± 7 years with a female predominance (sex ratio = 0.32). Three different molecules were involved: Acebutolol (n = 22), Atenolol (n = 2) and Propranolol (n = 1). The assumed average dose of ingested Acebutolol was 4000 [3200; 4800] mg. All patients were hospitalized in the first 3 h of taking and CS occurred within 3 h post ingestion in all cases. Bradycardia was reported in all patients who have ingested Atenolol or Propranolol. For patients who ingested Acebutolol, bradycardia was observed in 45% of cases (n = 10). Electrocardiographic signs included a ventricular atrioventricular block in 5 cases (as first degree in 3 cases and 3rd degree in 2 cases) and a membrane stabilizing effect in 10 cases. Digestive decontamination was performed in 14 patients (56%). All patients were treated using catecholamines: epinephrine in 18 patients (72%), both dobutamine and norepinephrine in 6 patients and only dobutamine in 3 patients. Glucagon was administered in 8 cases. Semi- molar bicarbonate intake as infusion of 500 ml into 30 min was prescribed to 10 patients who have membrane stabilizing effect. Eighteen patients required mechanical ventilation. The median duration of hospital stay was 72 (18; 120) h. Thirteen patients (52%) died within 24 h.

**Conclusion** Management of Cardiogenic shock complicating acute BB intoxication is largely symptomatic and based on vasopressors, positive inotropic agents. Sodium salts could be an alternative measure in cases of membrane stabilizing effect, and temporary cardiac electrical pacing if high degrees BAV. In refractory shock circulatory support should be initiated as early as possible.

**Competing interests** None.

### P104

**Purifying efficiency of CVVHDF and MARS in a simulated poisoning by verapamil**

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**Introduction** Drug poisoning is a frequent cause of hospital admission especially in intensive care unit (ICU). Despite progress, hospital mortality of severe acute poisoning admitted in ICU seems to increase. Purifying methods such as continuous haemodiafiltration veino-venous (CVVHDF) and molecular adsorbent recirculating system (MARS), were developed with promising clinical results (1). However no analytical study has quantified accurately their purifying efficiency. It has not been assessed efficiency of the different compartments of the MARS nor its advantages over other methods of dialysis and filtration. The objective of this study was to quantify the purifying efficiency of the different compartments of the CVVHDF and MARS and to compare their respective efficiency in an ex vivo model in the most favourable conditions for these methods to assess their capability maximum treatment.

**Materials and methods** We performed a ex vivo study based on manipulation bench simulating intoxication verapamil with a theoretical plasma doses of 1, 2.5 and 5 mg/l injected into a central compartment of 5 L. Sampling and assays were carried out at various points of the circuit CVVHDF and MARS. The EC extraction coefficient [EC = (in concentration – out concentration)/in concentration] was calculated for each compartment of the CVVHDF and MARS as well as the amounts withdrawn by the sum each compartment allows assessing the overall capacity of each technique. Three manipulations were performed for each concentration.

**Results** CVVHDF EC was 22, 22 and 28% for theoretical concentrations of 1, 2.5 and 5 mg/l. For the 3 previous concentrations, EC was not constant but decreased steadily over the sessions to cancel 0 ± 4 h for the different concentrations with even become negative after up to 14 h (end handling). At the end of handling the amount remaining in the central compartment was 6, 3 and 4% of injected dose. The amounts found in the effluent corresponded to 16, 20 and 28% of injected dose. In fact the amounts "not found" accounted for 78, 79 and 69% of injected dose. A desorption ultrasound membranes have revealed that verapamil was fixed by the membrane of the CVVHDF. During all handling (12–14 h) concentrations in the circuit output “patient” for the CVVHDF have never been undetectable but remained stable. The different compartments of MARS led instead to an undetectable concentration at the exit of "patient" circuit. Similarly the concentration of verapamil remaining in the central compartment was undetectable at the end of the handling and this for all three concentrations studied. The quantities withdrawn by the hemodiafiltration compartment MARS corresponded to 8, 18 and 12% of the injected quantity, EC varied between 10 and 20% remaining stable. The quantities withdrawn by the coal compartment activated MARS was 92, 82 and 89% of the injected amounts the EC induced by MARS cartridge was stable in the order of 70% throughout the manipulation.

**Discussion** CVVHDF and MARS are 2 effective purifying methods for removal of a simulated poisoning by verapamil. Their action and performance modes are very different. The membrane used in CVVHDF, adsorption verapamil realized 80% of the elimination of the toxic but with a residual salting out effect. A 8 h session of MARS had the effect of removing them undetectable concentrations in the central compartment. The purifying effect is bound to 80% to the effect of carbon cartridge and for 10–20% to hemodiafiltration.

**Conclusion** Clearance is a simple parameter that two determinants are EC and flow in the scrubber compartment. Pharmacokinetic modeling that require constant clearance. CVVHDF is characterized by a decrease in EC resulting in a variable clearance and preventing modeling. Elimination of verapamil is mainly due to unspecified binding properties. MARS is particularly effective in poisoning by verapamil. These results must be tempered by the fact that the conditions of our
manipulations, without protein in the central compartment, evaluate the maximum ex vivo treatment capacity.

**Competing interests**
None.

**P105**

**Acute beta blocker overdose management: Factors associated with cardiovascular mortality in a Caribbean intensive care unit**

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**Annals of Intensive Care 2017, 7(Suppl 1):P105**

**Introduction**
Beta-adrenergic antagonists are commonly used worldwide to treat hypertension, tremor, migraines, ischemic heart disease, heart failure, arrhythmias, portal hypertension, angina and panic attack. Propranolol, a beta-adrenergic antagonist with membrane stabilizing properties, is the most common toxic used in suicide attempts in Martinique.

Through respiratory depression, bronchospasm, bradycardia, severe hypotension, and seizures may result from beta blocker intoxication, cardiovascular depression appears to be the most common cause of morbidity and mortality in severe acute beta blocker poisoning. Massive beta-blocker ingestions may cause prolonged QRS intervals may also be associated with refractory cardiac failure. Our objectives were to determine factors that are associated with the development of cardiovascular mortality.

**Patients and methods**
We conducted a retrospective study over 10 years (January 2005 to December 2015), including all poisoned patients admitted and treated to the Emergency Department and the Intensive Care Unit. During this period, there were over 10 beta-adrenergic antagonist exposures per year reported by the medical records department. These poisonings accounted for an average of 5 deaths annually.

**Results**
Three hundred and eight patients (173 males/135 females) were admitted to the ICU for severe acute poisonings. Median age, 46.5% years (16–79); SAPS II, 120 (49–94). Among these 308 patients, 100 had ingested high doses of cardioxtoxins (class I anti-arrhythmic drugs (40%), β-blockers (15%), calcium-channel blockers (10%)). Fifty patients (50%) survived, including 18 to prolonged cardiac arrest. Bad prognostic factors in ECLS-treated poisoned patients for beta blocker poisoning, were as follows: QRS enlargement on admission, SAPS II score on admission, Extracorporeal Life Support (ECLS) performance under massage, potential co-ingestants, arterial pH and lactate concentration (10.5 mmol/l).

**Conclusion**
The most important factor associated with an increased risk of cardiovascular mortality in beta blocker poisoning is the exposure to a beta blocker with stabilizing activity. The identification of risk factors allows physicians to identify patients at greatest risk. ECLS appears to be an efficient salvage technique in case of refractory toxic cardiac failure or arrest.

**Competing interests**
None.

**Reference**
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**P106**

**Accidental poisoning in children aged under 6 years old admitted in pediatric intensive care unit: For which toxic?**

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**Introduction**
Accidental acute intoxications are part of pediatric medical emergencies of our hospital experience. Children are most at risk of accidental poisoning due to characteristics of their development and their environment. Our Goal is to point to the toxic substances involved and to draw up the outcome profile in these patients.

**Patients and methods**
Retrospective study, covering all poisoning cases occurred in children under 6 years old and admitted to PICU in the university hospital Oran. We specified the characteristics of children, the nature of the toxic, symptoms on admission, the therapeutic aspect, the evolutionary mode and length of hospital stay. The duration of our study was to 4 years, from 2011 to 2015.

**Results**
In those years, the ICU admitted 365 poisonings occurring in children. They represent 13% of all admissions and 71% (260 children) under 6 years old. The average age of addicts was 5.08 ± 1.36 years old with a male predominance.

All product types were involved in the poisoning. According to data, 72% caustic toxic (189 cases) followed by drug poisoning (in 28 cases) and it is basically a case of psychotropic and narcotic (01 case), and then in descending order are rodenticides poison (19 cases), organophosphorus insecticides like (11 cases), Hydrocarbon (7 cases) ingestion of plants (3 cases), making cosmetics (2 cases) and poisoning of industrial glue (01 cases). Upon arrival at the hospital, 68% of young children had gastrointestinal disorders, and 2.3% had neurological disorders (seizures, ataxia and abnormal movements), 1.1% were in respiratory distress.

After observation for most children, symptomatic treatment was introduced, 6 patients have been put under non invasive ventilation, intubation and mechanical ventilation was required in 01 patient for 16 days also put under Hemodynamic Support. The outcome was favorable with an average hospital stay of 28 h in survivors, and 01 death occurred following an ingestion of hydrocarbons.

**Discussion**
Children in pre-school age are often at risk of domestic accidents in relation to their age and environment. They ingest products wrongly deconditioned or transferred into current use containers. These by catch happen to be in the kitchen at a large household or in the bathroom. They also ingest drugs for adults and thus higher dosage: these drugs are too easily left to reach those children who are not suspected faculties to identify seize and swallow what they take to candy. A study in India positioned intoxication hydrocarbons 1st place (50.9%), followed by organophosphates (11.9%), medication and caustic identify (4.8%) cases each [1]. In Oran (Algeria) taking caustic remains at the level of poisoning in children. Overall mortality by poisoning is lower in children compared to adults. (The products in question are heterogeneous drugs, household products, plants, pesticides) We recorded 01 death secondary to acute respiratory distress syndrome following ingestion by siphoning gasoline.

**Conclusion**
At the end of our study, it appears that the acute poisoning in children is a very common event. However, developments remain favorable in most cases. This should not obscure the potential severity of the poisoning, nor lose sight of the necessary prophylactic measures. This prophylaxis should include education of the population, prevention of certain tragedies related to toxins or aberrant behavior, and triggering alerts to reduce morbidity and mortality related to poisoning.

Our ACCIPED laboratory medical university Oran for a future mission secondarly and tertiary prevention through continuous training for general practitioners, to respond to any request for advice on diagnosis and treatment of these accidents, assess risk through teledmedicine and through a website continuously 24 h/24 h.
Competing interests
None.

Reference
1. Hanouz J‑L, Lammens S, Tasle M, Lesage A, Gérard J‑L, Plaud B. Preoxygenation with spontaneous breathing or noninvasive positive pressure ventilation in the presence of calibrated leak: an experimental study in healthy volunteers. Fanny Le Gall1, Jean‑Luc Hanouz1, Hervé Normand1, Nicolas Terzi1

P107
Effectiveness of preoxygenation with spontaneous breathing or noninvasive positive pressure ventilation in the presence of calibrated leak: an experimental study in healthy volunteers Fanny Le Gall1, Jean‑Luc Hanouz1, Hervé Normand1, Nicolas Terzi1

Introduction Noninvasive ventilation has become the preoxygenation reference technique. In PreOx study [1], Hanouz et al. showed that the preoxygenation is faster with positive pressure ventilation with or without positive end expiratory pressure (PEEP) than spontaneous ventilation. However, during preoxygenation imperfect seal between the face mask and patient’s face can induce a leak decreasing its effectiveness. We assume that noninvasive ventilation could cancel the effect of the leak in non-obese subjects.

Materials and methods Experimental prospective study, randomised in cross-over of healthy volunteers from March to July 2016, in Caen university hospital. None of them were obese, smoked or had a chronic lung disease. Healthy volunteers conducted a preoxygenation with fraction of inspired oxygen at 100%: spontaneous ventilation (SV) and noninvasive ventilation (NIV, pressure support +6 cmH2O and PEEP +5 cmH2O) without and with calibrated leak (SV-leak and NIV-leak). From the volunteer mouth at respirator, subjects breathed through a mouthpiece (with a nose clip), mount catheter, antibacterial filter and From the volunteer mouth at respirator, subjects breathed through a mouthpiece (with a nose clip), mount catheter, antibacterial filter and end‑tidal of carbon dioxide were measured. A PreOx was deemed effective if FeO2 > 90%.

The breathing patterns of every subject seems to influence the existance of inspiratory leaks. Some volunteers had more inspiratory leaks than others, and experienced a longer length of preoxygenation than others. This is probably explained by a different instantaneous inspiratory flow [2]. These results hypothesize that individual preoxygenation is needed for each subject.

Conclusion Noninvasive ventilation allows effective preoxygenation despite the existence of leak.

Competing interests None.

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P108
Evaluation of bag‑valve‑mask ventilation in manikin studies: What are the current limitations? Abdo Khoury1, Fatimata Seydou Sall2, Alban De Luca3, Aurore Pugin3, Sebastien Pili‑Floury4, Lionel Pazart5, Gilles Capellier6

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Annals of Intensive Care 2017, 7(Suppl 1):P108

Introduction Manikin-based studies for evaluation of ventilation performance show high heterogeneity in the analysis and experimental methods used as we pointed out in previous studies. In this work, we aim to evaluate these potential limitations and propose a new analysis methodology to reliably assess ventilation performance.

Patients and methods One hundred forty healthcare providers were selected to ventilate a manikin with two adult self-inflating bags in random order. Ventilation parameters were analysed using different published analysis methods compared to ours.

Results Using different methods impact the evaluation of ventilation efficiency which ranges from 0% to 45.71%. Our new method proved to be relevant and showed that all professionals tend to hyperventilate and revealed a significant relationship between professional category, grip strength of the hand keeping the mask and ventilation performance (p = 0.0049 and p = 0.0297 respectively).

Discussion Using adequate analysis methods is crucial to avoid many biases. Extrapolations to humans still have to be taken with caution as many factors impact the evaluation of ventilation performance. Healthcare professionals tend to hyperventilate with current devices.

Conclusion We believe that problems related to manual ventilation efficiency could be prevented by implementing monitoring tools in order to give a direct feedback to healthcare professionals regarding ventilation efficiency and ventilatory parameter values.

Competing interests None.

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P109
Manual ventilation: to intubate or not to intubate the patient, it is not the question—a manikin study Fatimata Seydou Sall1, Alban De Luca2, Aurore Pugin2, Christelle Vidal1, Franck Leroux1, Lionel Pazart1, Gilles Capellier1, Abdo Khoury1

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Annals of Intensive Care 2017, 7(Suppl 1):P109

Introduction Basic and advanced airway management as Bag‑Valve‑Mask (BVM) and Endotracheal Intubation (ETI) are procedures used in prehospital resuscitation. In this work, we compare manual ventilation performance with both techniques and analyse their variability and also try to assess what factors may affect their performance.

Patients and methods One hundred forty healthcare providers were selected to ventilate an intubated and not-intubated manikin with two adult bags for 5 min each in a random order. Ventilation performance were analysed using a new analysis methodology described in a previous study which takes into account ventilation variability.

Discussion In this study we aimed to compare the efficiency of ventilation with two adult bags in manikins with and without an endotracheal tube. We observed a significant difference in ventilation efficiency between the two conditions. The main finding of our study is that the ventilation performance showed high heterogeneity in the analysis and experimental methods used as we pointed out in previous studies. In this work, we aim to evaluate these potential limitations and propose a new analysis methodology to reliably assess ventilation performance.

Conclusion The results of this study suggest that the selection of the analysis method can significantly impact the evaluation of ventilation performance. This highlights the importance of using appropriate analysis methods to ensure accurate and reliable results.
Results Five hundred sixty ventilation tests (280 for ventilation using a facemask and 280 for ventilation with an Endotracheal Tube (ETT)) were performed by the 140 healthcare professionals. Results show a significant difference between ventilation performance with a mask compared to ETT (p < 0.05) with more ventilation efficiency when healthcare professionals ventilate with an ETT than a mask (37 vs. 21 ventilation tests). Furthermore, a significant relationship is observed between participants’ professional category, the size of the hand squeezing the bag and manual ventilation performance (p < 0.05).

Discussion Healthcare professionals have performed 560 ventilation tests on an ETT and a facemask. Among them, only 58 (10.41%) were efficient and 502 (89.6%) are inadequate (of which 11.4% insufficient and 78.2% excessive). Whatever the kind of ventilation technique used, they are still struggling to perform manual ventilation efficiently according to international guidelines.

Conclusion The high ventilation performance failure observed in this study shows that “to intubate or not to intubate the patient” is not the question, but we should focus on how to improve efficiency of manual ventilation performed by healthcare professionals.

Competing interests
None.

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P110
A novel and global approach of ICU ventilator ergonomics
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Introduction Devices’ ergonomics are major determinants of task failure, especially in stressful environments. The aim of the study was to provide a global evaluation of ICU ventilator ergonomics using novel exploration tools.

Materials and methods Devices and physicians 6 new-generation ventilators were evaluated (Dräger Evita V300, Covidien PB980, Philips V680, Hamilton S1, GE CareScape R860, Maquet Servo-U) and compared to an old one (Carefusion Avea). 20 senior physicians were included, each testing up to 4 ventilators in a randomized order. Objective task completion 11 specific tasks were to be completed. The test was a failure if the correct response was given ≥120 s, or in case of incorrect response or abandoned task

User-friendliness evaluation System Usability Scale (SUS) was developed to measure device’s usability. It has a range of 0–100, the highest score being the best value. Mental Workload evaluation (MWL) is an exploration tools.

Physiological measurement Pupil diameter modifications were assessed by eye-tracking (SMI ETG 1); heart, respiratory rate and thoracic volume variations were measured with a dedicated device (Hexoskin).

Results No users could set inspiratory flow on Servo U, and only 18% succeeded with S1. Servo U had the worst global results (failure rate = 42%) and Avea the best (failure rate = 13%) (p = 0.12). Among all ventilators, Avea had the best SUS and TLX values, and Philips V680 the worst (p < 0.05). Eye tracking, respiratory rate and tidal volume activation differed between ventilators (p < 0.05). V300 caused the higher eye-tracking activation rate when compared to Avea (p = 0.03) and R860 (p = 0.019) (Fig. 27).

Conclusion Ergonomics evaluation is mandatory when evaluating new devices in the ICU. Most ICU ventilators presented poor HMI capabilities, thus allowing the occurrence of various hazards and failure.

Competing interests
None.

P111
Which of the last-generation ventilators is the most suitable for emergency transports and inter-hospital transfers?
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Introduction Hypotension and oxygen desaturation are the most common critical events occurring in 17% of emergency transports. They are mainly related to ventilator issues and the type of ventilator might mitigate the rate of adverse events. We decided to compare the most suitable ventilators for out-of-hospital transport.

Materials and methods This experimental bench test was designed to evaluate the technical characteristics, ergonomics, accuracy of volume and pressure delivery, triggering performance, pressurization and depressurization capacity, patient asynchrony and leak compensation of transport ventilators. We simulated many patient profiles and conditions by adjusting lung compliances (30, 100 ml cmH2O−1) and resistances (5, 20, 50 cmH2O L−1 s), inspiratory efforts (2.5, 5, 10 cmH2O), and leakage levels (<1, 3, 6, 12 L min−1). The ventilators were used in pressure control, volume control, and pressure support ventilatory modes in non-invasive and invasive settings.

Results Even if the technical characteristics of portable ventilators are quite similar, their ergonomics and performance are unequal. Major differences were found on tidal volume delivery, with mean relative errors ranging from +1.7 to −14.9%. Triggering delays and pressurization capacity also showed significant heterogeneity with a mean pressure time product (PTP) varying from 883 to 3018 cmH2O ms. The resistance of the expiratory circuit also differed across ventilators (from 1.7 to +1.4 cmH2O L−1 s), impacting the mean exhalation time and inducing air trapping and dynamic hyperinflation. Finally, while all ventilators were able to synchronize with patient’s demand at baseline (leakage <1 L min−1), only one showed adequate patient synchrony at high leakage level.

Discussion The main characteristics of last-generation ventilators are comparable, but major differences remain, especially in the conception of their basic ventilation modes. For instance, Hamilton T1 VC mode is actually a hidden pressure control mode that does not ensure
tidal volume accuracy under changing lung conditions, unlike specified by international guidelines.

**Conclusion** While it is still difficult to determine which ventilator best meets clinician expectations, this study will help to select the most appropriate device in regard to his own needs and make a clear comparison between ventilator performances.

**Competing interests** Abdo Khoury reports invitations to conferences from Resmed and Air Liquide.

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**P112**

**Variability of tidal volume in assisted mechanical ventilation in ARDS: a bench study**

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*Annals of Intensive Care* 2017, 7(Suppl 1):P112

**Introduction**

Even though limiting tidal volume (TV) in ARDS patients is recommended, this goal may not be achieved once spontaneous breathing comes up and assisted modes are used. Furthermore ICU ventilators offer numerous assisted ventilation modes that work differently across the brands. We underwent present bench study to systematically investigate the effect of assisted mechanical modes on a single ICU ventilator on size and variability of TV at different breathing frequencies (f), patient effort and ARDS severity.

**Materials and methods**

We performed a bench study in our university laboratory on an ICU ventilator (V500 Infinity, Dräger, Germany) using ASL 5000 lung model. Compliance was set at value mimicking mild, moderate and severe ARDS as recently reported. Thirteen assisted ventilation modes were tested falling into three categories, namely volume controlled ventilation with mandatory minute ventilation (VCV-MMV), pressure-controlled ventilation (PCV) including airway pressure release ventilation (APRV) and biphasic positive airway pressure (BPAP), and pressure support ventilation (PSV). fand effort were tested each at two levels for each ARDS severity in each mode. TV was expressed as median (first-third quartiles) and compared across modes using non-parametric tests. The probability for TV > 6 ml/kg ideal body weight (IBW) was assessed by binomial regression and expressed as odds ratio (OR) with 95% confidence intervals (CI). The variability of TV was measured from the coefficient of variation.

**Results**

The distribution of TV over all f, effort and ARDS categories significantly differed across modes (P < 0.001, Kruskal–Wallis test). TV was significantly greater with PSV (420 mL [332–527]) than with any other mode except for the three modes accommodating a variable PS level. The risk for TV to be greater than 6 ml/kg IBW was significantly increased with spontaneous breaths assisted by PSV modes (for PSV OR 19.36; [12.37–30.65]) and significantly reduced in APRV (OR 0.44; [0.26–0.72]) and PSV with guaranteed volume mode. The risk increased with increasing effort and decreasing f. Coefficients of variation of TV were greater for low f and for VCV-MMV and PCV modes. APRV had the greatest within-mode variability.

**Conclusion** The ventilation mode had an important impact on TV in this study. The risk of TV > 6 ml/kg IBW was significantly reduced in APRV and PSV with guaranteed volume mode. APRV had the highest variability. PSV with guaranteed volume could be tested in ARDS patients.

**Competing interests** None.

**P113**

**Reusable versus single use fibroscope in the ICU: a medico-economical evaluation in the ICU**

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*Annals of Intensive Care* 2017, 7(Suppl 1):P113

**Introduction** Single-Use Fibscopes (SUF) have recently become available for ICU clinical practice to overcome 3 main problems associated with Re-Usable Fibscopes (RUF) resulting in systematic: immediate availability of a sterile device, short lag time to endoscopic treatment and removal of the cross contamination risk. However endoscopy of the upper airway and lung (EAWL) is a medical act that generates significant costs. We thus conducted a medico-economical evaluation of RUF versus SUF.

**Patients and methods**

The study was conducted over 1 year (2015) in 3 French university hospitals. The medical evaluation was conducted in one of them while the cost analysis included the 3 hospitals.

**Discussion**

SUF is efficient and performant for EAWL. Although costs associated with the use of SUF and RUF for ICU EAWL are close, selecting SUF induces substantial savings. The permanent and immediate availability of sterile equipment allowing to significantly shorten the lag time to treatment may have important clinical outcome for the patients.

**Conclusion** At a time when health authorities are focusing their efforts on the prevention of nosocomial infections, elimination of endoscopes-related cross-contamination risk by using SUF, especially in ICU is a major issue. Improvement of Suction capability will promote clinical superiority of SUF over RUF.

**Competing interests** Dhonneur Gilles is consultant for Ambu.

**P114**

**Nociception resulting from oro-tracheal intubation: an observational study**

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**Introduction** Oro-tracheal Intubation (OTI) is thought to be one of the highest noxious stimulation for the patient but this dogma has never been clearly proved. We tightly monitored hemodynamics (HD), noceception, and consciousness during OTI with the aim of pragmatically revisiting this dogma.

**Patients and methods**

After ethic comity approval, consenting informed patients scheduled for elective cardiac or vascular surgery were included in this observational prospective study. Patients were orally premedicated (hydroxyzine) 1 h before admission into the
operating room, where invasive blood pressure (BP), Bis-spectral (BIS) and Analgesia Nociception Index (ANI) monitoring were installed. The induction procedure of anesthesia with target-controlled infusion was standardized using sufentanil and propofol. Timed OTI maneuver was performed after monitored (corrugator supercilii muscle) deep neuromuscular blockade (atracurium) was installed. OTI maneuver which duration could not exceed 30 s was truncated in 3 periods (P) of predefined duration. P1 lasted 10 to 15 s and corresponded with direct laryngoscopy. P2 lasted 5 to 10 s and corresponded with tracheal tube (TT) manipulation. Finally, P3 coincided with TT cuff inflation to control 35 cm H2O lasted less than 5 s. Recorded HD, BIS and ANI data were analyzed at 6 predefined time points. T0: stable baseline values after monitors installation, T1: post induction of anesthesia and just before OTI, T2: end of P1, T3: end of P2, T4: end of P3, T5 and T6, respectively 1 and 5 min after the end of OTI. Recorded parameters evolution was compared using standard statistics.

Results Thirty-nine patients were included in this trial, but 4 were secondarily excluded because of OTI lasting more than 30 s. Figure 28 shows mean (SD) variations of monitored parameters: mean arterial pressure (MAP), heart rate (HR), BIS, and ANI, from T0 to T6. P1 is characterized by a remarkable stability of all measured parameters. P2 and even more P3, are associated with intense and significant HD and ANI variations. ANI variations are completely superimposed, but opposite, to those of MAP and HR. BIS variations during TI were negligible.

Conclusion Our data does not support the dogma but rather suggests that laryngoscopy may not be the most intense nociceptive stimulus applied to patients during OTI. Both laryngeal and/or tracheal stimulation resulting from tracheal tube passage and cuff inflation seem to promote very intense nociception. Interestingly, such phenomenon should be erased by either laryngeal mucous topicalization or superior laryngeal nerve block.

Competing interests None.

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P117
Comparisons and time course of mixed venous and central venous saturation are independent of severe sepsis and septic shock origin
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Introduction Low (<70%) mixed venous saturation (SvO2) was one of the independent variable related to mortality in septic shock patients [1]. Nevertheless, values of central venous saturation (ScvO2) may be considered as a surrogate to SvO2 for early goal directed therapy in severe sepsis and septic shock. In sepsis, difference between ScvO2 and SvO2 (ΔSvO2) was approximately of 5% [2]. This delta SvO2 ignores the origin of sepsis: supra or infra-diaphragmatic where this gradient could be higher due to increased oxygen extraction by organs under diaphragm (gut, liver and kidney). We compared the correlation between ScvO2 and SvO2, the time course of delta SvO2 and effects of therapeutic challenge on delta SvO2 between patients in severe sepsis or septic shock due to supra or infra-diaphragmatic infection. Enter the text.

Patients and methods We included patients in severe sepsis or septic shock monitored by Swan Ganz catheter and continuous ScvO2, hospitalized in a 24 bed medico-surgical ICU of CHU-Charleroi. Hemodynamic data and delta SvO2 were recorded from catheter insertion to withdrawal and before and after each therapeutic intervention (fluid challenge, dobutamine initiation or dosage modifications). Demographic data and mortality were also collected. Graphpad prism was used for statistical analysis and graph, Bland–Altman analysis was used and results were compared by Mann–Whitney U test when not specified. Data were expressed in median values (IQR 25–75 percentiles).

Results 34 patients were included which 26 presented a septic shock of upper diaphragmatic origin. Median Apache II score was 24 (16–30). Eighteen patients (53%) out of 34 died before hospital discharge. At inclusion, mean arterial pressure, SvO2, cardiac index and norepinephrine infusion rate were respectively 71 (67–82) mmHg, 66 (59–72%)% 2.7 (2.2–3.4) L/min/m2, and 0.09 (0.00–0.50) (mcg/kg/min).

For all patients included, a total of 442 paired blood samples were obtained (321 for upper diaphragmatic group). ScvO2 overestimate SvO2 by an average of 4.96% (−6.4 to 16.3) for all paired data. No difference between upper (bias 5.24, limits −6.1 to 16.56) and under diaphragmatic (bias 4.21, limits −7.1 to 15.33) paired (ΔSvO2) was observed (p = 0.1).

Time course of ScvO2 and SvO2 were similar during the study period and independent of the sepsis origin.

No difference in ΔSvO2 was observed in relation the baseline SvO2 values (<50% and higher values (p = 0.11)).

Median SvO2 and ScvO2 were not statistically different before (64 [55–72] and 70 [63–78]% respectively) and after intervention (67 [56–75] and 72 [65–78]% respectively (p = 0.27 for SvO2, and p = 0.40 for ScvO2). Therefore, ΔSvO2 was the same before and after interventions in the 2 groups of patients (p = 0.09) (Fig. 29).

Conclusion As in other studies, ScvO2 overestimate SvO2 by nearly 5.0% in patients with septic shock independently of the sepsis origin. ScvO2 could be used as a surrogate of SVO2 in the management of septic patients.

Competing interests None.

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2. Brocc O, Reinhart K. Intens Care Med. 2005; 31: 911–13.
Introduction To investigate whether respiratory variations of the inferior vena cava diameter (iVCD) assessed during a simple standardized breathing maneuver predict fluid responsiveness in spontaneously breathing patients with septic acute circulatory failure and irregular heartbeats.

Patients and methods This prospective, bicentric study, was performed in the intensive care units of a general and a teaching hospital. Spontaneously breathing patients with cardiac arrhythmia considered for volume expansion for clinical signs of acute circulatory failure related to sepsis were included. Echocardiography and Doppler ultrasound were used to record the stroke volume (SV) and IVC collapsibility index (iVIc) defined as [maximum expiratory diameter (edIVC) − minimum inspiratory diameter (idIVC)]/edIVC × 100 at baseline and after a 500 ml colloid infusion. Vena cava pertinent diameters were measured 15–20 mm caudal to the hepatic vein junction and recorded by bi-dimensional imaging on a subcostal long axis view. We measured the edIVC and idIVC during standardized (st) and unstandardized (ns) breathing, and calculated cIVCst and cIVCns before fluid loading. Standardized respiratory cycles consisted of a deep standardized inspiration followed by passive exhalation. A positive response to fluid loading was defined as an increase of the SV > 10%.

Results Fifty-five patients were included, 29 (53%) with atrial fibrillation and 25 with atrial Extrasys-t >6/min. Twenty-nine (53%) patients were responders to fluid loading. A cIVCns > 19% predicts fluid responsiveness with a sensibility of 83%, a specificity of 68%, and an area under the ROC curve of 0.83 (95% CI 0.72–0.94). A cIVCst > 39% predicts fluid responsiveness with a sensibility of 93%, a specificity of 88%, and an area under the ROC curve of 0.93 (95% CI 0.86–1). The area of uncertainty ranged between 10 and 13 mm.

Discussion A simple standardized inspiratory maneuver improves the accuracy of cVIc to predict fluid responsiveness with high specificity and sensibility and a limited grey area in patients with cardiac arrhythmia. Moreover, idIVCst demonstrates very good performance to predict fluid responsiveness. This easy to acquire parameter may be more suitable for bedside patient’s management.

Conclusion cIVCst is a powerful index to predict fluid responsiveness in spontaneously breathing patients with sepsis and irregular heartbeats. Interestingly, idIVCst shows similar performance to predict fluid responsiveness with a much easier assessment, improving feasibility in clinical routine.

Competing interests None.

Discussion Several studies in emergency and pneumology departments have investigated global hemodynamic parameters to predict outcome of PE patients. Low systolic blood pressure called “shock” and right ventricular dysfunction were identified as predictors of bad outcome. We were interested in the exploration of tissue perfusion because in septic shock several studies highlighted the discordance between global hemodynamic parameters and tissue perfusion. Moreover, during severe infections, hypoperfusion markers such as lactatemia, mottling or oliguria have been identified as predictive of ICU mortality. Recently, Vanni et al. [2] suggested lactate could be useful to identify normotensive PE patients at high risk of early death.
Here, we found that, at admission, non-survivors patients at day-28 had more severe tissue hypoperfusion than non-survivors. In addition, mortality increased with the number of tissue hypoperfusion parameters. These results have to be confirmed prospectively in a large cohort of PE patients but suggest that tissue hypoperfusion could be helpful for triage and potentially for guiding treatment.

**Conclusion** In a retrospective study, we observed an association between tissue hypoperfusion parameters and 28-day outcome in ICU patients admitted for acute symptomatic PE.

**Competing interests** None.

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**P122**

**PEEP sparing in prone position has a weak but significant positive effect on cardiac index in ARDS patients**

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**Annals of Intensive Care 2017, 7(Suppl 1):P122**

**Introduction** The PROSEVA study was the first randomized controlled study having shown a positive effect of prone position (PP) on ARDS mortality [1]. Unlike previous randomized controlled trial, PEEP was adjusted with a PEEP-FIO2 table, whose consequence was a small but significant decrease in the PEEP level. The aim of this study was to investigate the effect of PEEP variation during prone position on cardiac index in ARDS patients.

**Patients and methods**

Single center retrospective observational study performed on ARDS patients hospitalized in a medical ICU between July 2012 and March 2016. Patients included were adults fulfilling the Berlin definition for ARDS, undergoing at least one prone position session, under hemodynamic monitoring by the Picco device, with availability of hemodynamic measurements performed before (T1), at the end (T2), and after the prone position session (T3). Prone position sessions were excluded if they were performed >7 days after ARDS onset. The following variables were recorded: demographic, SAPSII, ARDS severity and risk factor, SOFA score and cumulative fluid balance at PP onset, delay between ARDS session and PP session, hemodynamic, arterial blood gas, ventilatory settings, plateau pressure, catecholamine dose and additional treatments. Statistical analyses were performed using prone position session as statistical unit and mixed models taking into account both multiple prone position sessions by patient and multiple measurements during a prone position session. Variables associated with cardiac index with a p value below 0.1 in univariate analysis were selected for inclusion in a multivariable mixed logistic regression model, using backward stepwise descending selection. p < 0.05 was chosen for statistical significance. Data are expressed as mean ± standard deviation.

**Results**

85 patients fulfilled the inclusion criteria over the study period, totalizing 149 prone position sessions (2 ± 1 sessions per patient). PEEP level decreased significantly from 9 ± 3 to 8 ± 2 cm H2O between T1 and T2. PEEP decreased by at least 5 cm of H2O in 18 (12%) of the PP sessions. Multivariate analysis identified 7 variables independently associated with cardiac index (model 1, cf. Table 2). Multivariate analysis also identified measurement time, age, sex, PEEP and pH as being independently related to global end-diastolic volume. After removing the effect of global end-diastolic volume by forcing it out of the model, PEEP became significantly related to cardiac index (model 2, cf. Table 22), with a coefficient B suggesting a marginal effect on cardiac index (−0.03 l min⁻¹ m⁻² per 1 cm H2O increase of PEEP).

**Conclusion** PEEP decrease during prone position as a consequence of improvement of oxygenation has a marginal but significant positive effect on cardiac index, related to an increase in global end-diastolic volume and probably venous return.

**Competing interests** None.
Table 22 Multivariate analyses of variables associated with cardiac index

|                     | Model 1 β | p     | Model 1 β | p     | Model 2 β | p     | Model 2 β | p     |
|---------------------|-----------|-------|-----------|-------|-----------|-------|-----------|-------|
| Age                 | −0.043    | <0.001| −0.021    | <0.05 | −0.021    | <0.05 | −0.023    | <0.01 |
| SOFA score          | 0.071     | <0.001| 0.103     | <0.001| 0.001     |       | 0.013     | <0.05 |
| Neuromuscular blocking agents (ref = no) | 0.450     | <0.001| 0.336     | <0.05 | 0.336     | <0.05 | 0.336     | <0.05 |
| Global end-diastolic volume (mL m⁻²) | 0.003     | <0.001| −0.232    |       | −0.232    |       | 0.033     | <0.05 |
| Dobutamine dose (µg kg min⁻¹) | −0.021    | <0.01 | −0.232    |       | −0.232    |       | 0.033     | <0.05 |
| PaCO₂ (mmHg)        | 0.014     | <0.01 | 0.013     | <0.05 | 0.013     | <0.05 | 0.013     | <0.05 |
| PaO₂/FiO₂ (mmHg)    | −0.002    | <0.001| −0.001    | <0.05 | −0.001    | <0.05 | −0.001    | <0.05 |
| PEEP (cm H₂O)       | NS        | NS    | −0.330    | <0.05 | −0.330    | <0.05 | −0.330    | <0.05 |
| Time between ARDS onset and PP session | NS        | NS    | −0.044    | <0.05 | −0.044    | <0.05 | −0.044    | <0.05 |
| Renal replacement therapy (ref = no) | NS        | NS    | −0.355    | <0.05 | −0.355    | <0.05 | −0.355    | <0.05 |
| Measurement time [ref = before PP (T1)]    | NS        | NS    | <0.001    |       | <0.001    |       | <0.001    |       |
| End of PP (T2)      | 0         |       |           |       |           |       |           |       |
| After PP (T3)       | −0.204    |       |           |       |           |       |           |       |

P123

Impact of implementing a sodium phenyl-acetate and sodium benzoate delivery protocol on hyperammonaemia

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Introduction

Hyperammonaemia is defined as ammonia above 80 µmol/L when child is under 1 month old and 55 µmol/L when over 1 month old. Hyperammonaemia over 200 µmol/L was proved to be an independent factor of mortality. The recommended treatment for hyperammonaemia due to inherited error of metabolism is based on the association of sodium phenyl-acetate and sodium benzoate called Ammonul®. In order to improve management of hyperammonaemia in pediatric intensive care unit (PICU), an Ammonul® delivery protocol was implemented since the 30th August 2008. The purpose of our study was to evaluate the impact of this delivery protocol on morbidity and mortality for patient admitted in PICU for hyperammonaemia.

Patients and methods

A retrospective study was conducted from January 1st 2000 to May 31st 2016 in the PICU of the CHU Sainte-Justine, Montreal, Canada. Patients were aged 9 months to 452 months. Forty-two patients were studied, 8 in the without protocol group and 14 in the with protocol group. Patients were aged 9 months to 452 months in the without group and 228 to 512 months in the with group (p = 0.09). There was no difference between groups in term of weight (6 [3–43] vs 24 [3–67]), ammonia level (328 [75–2022] vs 239 [150–830]) and severity score (PIIMII 18 [5–54] vs 19 [2–100]). The principal cause of hyperammonaemia was inherited error of metabolism in both groups (6 (75%) vs 9 (64%)). The median delay between diagnosis and Ammonul prescription was 100 min [75–209] in the without group and 47 min [0–81] in the with group (p < 0.001), the median delay between diagnosis and Ammonul administration was 173 min [95–339] in the without group and 173 min [73–276] in the with group (p = 0.63).

Conclusion

Our study showed that the implementation of a sodium phenyl-acetate and sodium benzoate delivery protocol shorten the delay between diagnosis of hyperammonaemia and treatment prescription but failed to find any improvement in morbi-mortality.

Competing interests
None.

P124

Patients characteristic of French pediatric intermediate care units

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Introduction

In France, since 2006, legislation for pediatric critical care organization approved the establishment of pediatric intermediate care units, between pediatric wards and pediatric intensive care units. In pediatric intermediate units, children require close monitoring and/or continuous monitoring due to potential organ failure, without requiring resuscitation. The aim of this study is to describe patients characteristics of French pediatric intermediate care units.

Patients and methods

We performed a prospective observational cohort study in pediatric intermediate care units of seven regional hospitals in northern of France. All consecutive patients under 18 years, admitted in these seven pediatric intermediate care units were included (September 2012–January 2014).

Results

Among 2909 consecutive screened patients, 2868 patients were included. Sex ratio was 1.26. Median of age was 29[5–103] months. Median of length of stay was 1 [1–3] days. Thirty-three percent had comorbidities. Seventy per cent were transferred from emergency unit. The type of stay was medical for 95%. The primary reason for admission was essentially respiratory (44%) and neurologic (22%). Infection was the main cause of respiratory failure (79%). The destination was pediatric ward in 53% and home in 36%. Three per cent were transferred in pediatric intensive care unit and one patient deceased.

Conclusion

The patients hospitalized in pediatric intermediate care unit were young. The length of stay in these units was short and the respiratory failure was more frequent.

Competing interests
None.
Validation of three pediatric early warning scores in seven French pediatric intensive care units

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Introduction Pediatric Early warning systems were created to quantify severity of illness across in hospitalized children in pediatric ward or emergency unit. Pediatric intermediate care units are alternative structures for moderate ill children. The aim of this study was to assess the validity of three pediatric early warning scores in pediatric intermediate care units.

Patients and methods We did a prospective, observational, multicenter cohort study in seven French regional hospitals (09/2012–01/2014). All consecutive children <18 years were included. Three scores (PAWS, PEWS, and BedPEWS) were calculated each 8 h and more if deterioration. Binary outcome criteria were "medical call"; and "admission to Pediatric intensive care Unit (PICU)". When one or two monitoring parameters necessary to calculate the score were missing, the score was still calculated and the missing value was considered normal and these scores were called “imputed scores”. We used areas under the curve to estimate discrimination.

Results 2868 children were included for a total of 19,071 observations for calculating the three scores. Median scores for the three scores were respectively 2, 2 and 1. Medical call was observed in 11% (n = 2056), and admission to PICU in 0.45% (n = 85). AUCs calculated for the three scores for predicting medical call were ranged from 0.92 to 0.93. AUCs for predicting PICU admission were ranged from 0.89 to 0.92 (Table 23).

Conclusion The three Pediatric Early warning scores, developed from pediatric ward and emergency department, can be used to detect deterioration requiring a medical intervention or PICU admission in hospitalized children in pediatric intermediate care unit.

Competing interests None.

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Breastfeeding disruption during hospitalisation for acute bronchiolitis in children: determining factors in a French teaching hospital

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Introduction No study have been published to specifically evaluate or measure breastfeeding disruption during hospitalisation for bronchiolitis although the high number of hospitalisation for bronchiolitis in children potentially breastfed makes it an important health issue. We conducted a pilot study to try to measure that risk.

Patients and methods This is a single center prospective observational study conducted between the 1st of October 2015 and the 15th of February 2016 in a tertiary care teaching hospital. All patients under 6 month hospitalized with acute bronchiolitis and receiving at least partial breastfeeding were considered for study. Patients discharged at home whose parents accepted to be contacted by phone were prospectively included. The primary outcome was unwanted weaning from breastfeeding. The secondary outcome was trying to identify risk factors for weaning. Data are expressed as median values (with minimum and maximum value) for continuous variables, and number and/or frequency (%) for binary or categorical data.

Results During the study period 144 breastfed patients under 6 month were hospitalised at our hospital for bronchiolitis, 84 patients (58%) could be included in the study. Length of hospital stay was 3 days [1; 34] and 27 patients (32%) spent some time in PICU. Length of hospital stay was 3 days [1; 7] and 34 patients (40%) received standard oxygen for 3 days [1; 9], 34 patients (40%) received no respiratory support. Forty-five (53%) patients received nutritional support, either enterally (N = 38, 45%) or parenterally (N = 5, 6%) or both (N = 2, 2%). Most breastfeeding mothers (96%) did not smoke and lived as a couple (98%), 2% of mothers were working at the time of hospitalisation. Sixty-five patients (77%) were exclusively breastfed before hospitalisation. The median delay for phone contact was 3 month [0.5; 6]. Forty-three mothers (51%) stated that their breastfeeding had been modified by the hospitalisation of their child: 17 stopped breastfeeding (Group 1), 12 switched from total to partial breastfeeding (Group 2) and 14 reduced breastfeeding without stopping (Group 3). Remaining mothers (Group 4) (41, 49%) stated to have kept breastfeeding as before or that their breastfeeding modification was personal choice. Patients whose breastfeeding was more impacted had a tendency to being younger with a median age of 34 days [3;166] vs 50 days [16;159] on admission, p = 0.06 and to be exclusively breastfed before (81 vs 73%, NS). No difference was observed for birth weight, gestational age, daily growth before stay, length of stay, need for respiratory support, need for nutritional support, need for PICU admission between groups 1 + 2 + 3 and group 4. Mothers stated first cause of breastfeeding disturbance had been lack of support and advices (32.5%) followed with some logistics problem (difficulties to draw breast milk, room accommodation, introduction of formula milk) in 15.7% cases and severity of child’s respiratory disease in 16.8% cases.

Discussion This is the first study to highlight the effect of hospitalisation for bronchiolitis on breastfeeding, and to question the various factors involved. Our study is subject to possible bias (single centered, long delay between hospitalisation and questionnaire) but we...
still achieved to reach a big number of patients (84 patients in one epidemic season). We observed a very high rate of breastfeeding disturbance during hospitalisation for bronchiolitis in our hospital (over 50%). We expected to find severity of respiratory disease as the first risk factor but only 16.8% of mothers pointed out severity of disease as a reason for unwanted weaning and respiratory distress on medical charts was also not significantly different. Lack of support from caregivers was the first factor pointed out. It is a modifiable factor for further practice, especially in younger infants for which breastfeeding has just begun and is still precarious.

**Conclusion** Bronchiolitis is a high risk event for breastfeeding with about half of mothers in our hospital either stopping or diminishing their breastfeeding during hospitalisation. Correct advice and support could be a determining factor of breastfeeding’s continuation and further studies should focus on interventions to prevent unwanted weaning.

**Competing interests** None.

P127

Factors associated with severe bronchiolitis in PICU: a retrospective study

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**Introduction** The explosion of non invasive ventilation (NIV) and high frequency canula therapy (HFNC) in bronchiolitis for the last 10 years led to a decreased rate of intubation [1]. There are lacking data concerning the clinical and biological factors associated with severe bronchiolitis defined by the need for an intensive level of NIV or for invasive ventilation (IV).

**Patients and methods** Two hundred and fifty-two patients aged of 6 months and 6 years were included in a retrospective study conducted over a three winter period from 2013 to 2016 in a 16 bed French PICU. We aimed to compare patients hospitalized for severe bronchiolitis, defined by the need for an intensive level of NIV or for invasive ventilation (IV) to non severe patients that were treated with nasal continuous positive airway pressure (nCPAP) or HFNC. We secondly analyzed the evolution of patients characteristics over a three winter period and factors associated with HFNC failure. Clinical and biological factors were recorded at the admission of the patients and 4 h after (H4).

**Results** One hundred patients were included in the non severe bronchiolitis group and 152 in the severe group. Factors significantly associated with severe bronchiolitis were a young age, a small weight and the presence of apneas at the admission. The Wang severity score, the heart rate, the apH and the PaCO2 were significantly different between the two groups at the admission and at H4. The level of oxygenation at H4 were also higher in the non severe group. The rate of bacterial coinfection, the length of stay in PICU and of ventilation were higher in the severe group. Over the 3 years period, the rate of IV and NIV bilevel positive airway pressure significantly decreased in parallel with the increased utilization of HFNC. Comparing patients responders or not to HFNC, the weight, the Wang score, the respiratory rate, the apH and the PaCO2 at H4 were significantly different.

**Discussion** First, we found that clinical and biological factors associated with the need for a NIV bilevel positive airway pressure were similar of those associated with the need for IV previously described [2]. We can hypothesize that patients at risk of severe bronchiolitis and NIV failure are actually better identified with prompt modifications of ventilation parameters and a greater tolerance for permissive hypercapnia under NIV support. All of these factors lead to a reduction of the rate of intubation [1]. Secondly, patients hospitalized seemed to be less severe according to biological parameters (apH and PaCO2) and the decreased need for IV all along the three seasons studied. This could be in relation with an earlier recognition of critically illness patients and no delayed respiratory support treatment. This could also be related to an increased administration of HFNC to patients usually treated with standard oxygen treatment. Finally, we reported factors associated with HFNC failure. Even if HFNC is not actually recommended in the first line treatment of bronchiolitis, some authors described its efficiency in this pathology, with similar results as nCPAP and superiority to standard nasal oxygen with reduced respiratory rate, respiratory work, length of stay and level of oxygenation.

**Conclusion** NIV is a useful respiratory support technique in paediatric patients with bronchiolitis. Younger and heavier patients, clinical severity score, respiratory and heart rate or biological parameters like apH and PaCO2 are factors associated with an intensive level of NIV.

**Competing interests** None.

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P128

Practice survey on prone positioning in French-speaking pediatric intensive care units (nursing part)

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**Introduction** The prone positioning (PP) is a strategy widely used in the treatment of severe forms of acute respiratory distress syndrome (ARDS) in adults. Its early use significantly reduces mortality [1]. However, the studies do not strongly demonstrate its prognostic impact in pediatric ARDS. The aim of this study was to describe the prone positioning practices in the French-speaking pediatric intensive care units (PICU).

**Patients and methods** This survey was conducted by email questionnaire to pediatric intensivists belonging to the French Society of Intensive Care Medicine and the French-speaking Group of Pediatric Intensive Care and Emergency Medicine. It was conducted from February to May 2016. The survey was addressed to doctors, nurses, physiotherapists practicing in PICU. It included 29 questions about indications, contraindications, techniques and medical devices used, and complications.

**Results** One hundred and three persons answered (69 doctors and 33 nurses) which work in 28 French hospitals and 1 Canadian hospital. The thoraco-abdominal support is use whatever the age (neonates, infants, children) and frequently made with non-specific material: sheets and undersheets (64%), cocoons with balls (44%), pillows and bolsters (38%). The members are placed in flexion or half-flexion (71% of interviewed persons) without systematic changes of the positions. The PP is usually treated during the dressings (72% of interviewed persons), less frequently to achieve a toilet (44%), radiography (43%) or physiotherapy session (29%). Seventy-six percent of interviewed persons say that the kind and location of prostheses (catheter, drains, stomy) are not contraindications to the PP. Concerning respiratory care 74% of interviewed persons frequently use a closed system for tracheal aspirations. Moreover, 45% of interviewed persons say that respiratory physiotherapy sessions are not classically realized in PP periods. Concerning the enteral nutrition 59% of persons do not change their practices.
during PP and only 10% reduce or stop it. Finally the mucouscutaneous complications are the most frequent (54% of interviewed persons) just before the displacement of protheses (37%). The prevention of cutaneous pressure ulcers by the use of specific mattress is widespread (61% of interviewed persons) but no supplemental care in prone compared to supine positioning.

**Discussion** The survey aimed to describe the techniques and modalities of PP in children, as well as several aspects of care associated with PP. The results show that the quality of care seems unaffected by the PP. The interviewed persons do not reported technical contraindication of PP. However, a special attention should be put on hemodynamic instability, tracheostomy or umbilical venous catheter. The survey reported the same complications in children and adult [2]. Finally, we also report the absence of paramedical care protocol for the PP. Future clinical studies will assess the impact of nursing protocols to limit complications, improve quality and comfort of care in children during PP periods.

**Conclusion** The prone positioning in French-speaking pediatric intensive care units is not very well protocolized. Nevertheless, the care not seems to be significantly impacted by the prone positioning. Future research should focus on the evaluation of prone positioning effects on patient care.

**Competing interests** None.

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**P129**

**Implementation of a weaning protocol of mechanical ventilation in a pediatric intensive care unit in Oran, Algeria**

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**Annals of Intensive Care 2017, 7(Suppl 1):P129**

**Introduction** The Mechanical ventilation (MV) is the main technical motivating ICU admission and involves on average 30% of children (range 20–60%), half is extubated within 48 h. Weaning from mechanical ventilation, should be as early as possible in order to reduce the risk of iatrogenic complications, hospital stay in intensive care and costs. Our purpose is the implementation of a written protocol of weaning from mechanical ventilation in a pediatric intensive care unit.

**Patients and methods** This is a prospective study including all children aged between 1 month and 15 years, admitted to the pediatric intensive care unit at Canastel Hospital in Oran Algeria during the period from 1 March 2015 to 31 August 2016. The weaning protocol involves the systematic and daily search for 7 criteria for sevrabilité allowing extubation:

- Resolution of the case
- FiO₂ < 50% or SaO₂ > 90% and ≤5 cm H₂O PEEP
- Dobutamine or noradrenaline ≤ 0.2μg/kg/min
- Conscious or slightly sedated or Glasgow >12
- Cough, effective deglutition
- No surgery ≤12 h
- T < 38.5°C

**Results** We included 43 children. The average age is 6 years (1 month–14 years), sex ratio is 1.2. The most common reason for admission in ICU was the severe brain injury in 21.4% and the status epilepticus in 19%. The patients were intubated in 70.7% of cases for an alteration of the state of consciousness. Ten patients (25.6%) had a post extubation stay was 1 required reintubation within 24 h. A corticosteroids administered in 67.4% of patients in the 12 h before extubation. No deaths during the study.

**Conclusion** Weaning from mechanical ventilation is a key direction for the prognosis of ICU patients where the value of a weaning protocol with general and respiratory criteria to facilitate extubation.

**Competing interests** None.

**P130**

**Severe atypical pneumonia in critically ill patients: a retrospective multicentre study**

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**Annals of Intensive Care 2017, 7(Suppl 1):P130**

**Introduction** Chlamydia pneumoniae (CP) and Mycoplasma pneu- moniae (MP) are intracellular pathogens that account up to 20% of community acquired. These patients with “atypical pneumonia” may require ICU admission for acute respiratory failure.

**Patients and methods** Medical charts of adult patients hospitalized between 2000 and 2015 in the ICU of 20 French hospitals with proven atypical pneumonia (positive serology or PCR) were retrospectively reviewed. Patients with Mycoplasma pneumoniae infections were compared to microbiologically documented Streptococcus pneumoniae patients (SP).

**Results** 104 patients were included (71 men, 33 women) with a median age of 56 (44–67) years, mainly admitted to the ICU for acute respiratory failure (n = 96, 92%). Mycoplasma pneumoniae was the causative agent in 76 patients (73%) and Chlamydia pneumoniae in 28 (27%). Atypical pneumonia was more frequent during autumn and winter. Superinfection was reported in 19 cases (viruses in 47%). Clinically, 33 patients (32%) had at least one extra pulmonary symptom. Chest-X-ray disclosed an involvement of 2 [1–4] quadrants, alveolar opacities (n = 61, 75%), interstitial opacities (n = 32, 40%) or pleural effusion (n = 6, 7%). During ICU stay, 75 patients (72%) required mechanical ventilation. Among them, 34 had ARDS. More than one-third of the patients (n = 41) received vasopressors. ICU length of stay was 14 days [6–21] in patients that were discharged alive and 23 [18–41] in patients who died in the ICU-days; 11 (11%) patients died in the ICU. Factors associated with mortality, in univariate analysis, were age ≥65 years (p = 0.033), signs of respiratory distress (p = 0.017) and interstitial opacities on chest-X-ray (p = 0.017).
Characteristics at admission for SP and MP patients were compared (Table 24). HIV infection was more frequently associated with SP pneumonia (16 vs. 3%, \( p = 0.009 \)). SP patients presented more frequently with shock in (32 vs. 8%, \( p = 0.0004 \)) and acute renal failure (13 vs. 9%, \( p = 0.008 \)). SP pneumonia was associated with higher ICU mortality (20 vs. 5% at 28 days, \( p = 0.005 \)).

**Conclusion** Critically ill patients with severe atypical bacterial pneumonia have a 11% case fatality. This study maintains a high level of suspicion towards atypical pneumonia as compared to SP pneumonia.

**Competing interests**
None.

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**P131**

Severe pneumonia due to legionella pneumophilia: clinical characteristics and outcomes

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**Introduction** Legionnaire’s disease (LD) is a severe pneumonia commonly caused by *Legionella pneumophila* serogroup 1. Despite advances in critical care management and antimicrobial therapy its morbimortality remains still high. No epidemiologic studies had been reported in Tunisia.

**Objectives** To study characteristics and outcomes of patients admitted in ICU with severe community acquired pneumonia due to LD over 17 years.

**Patients and methods** It was a retrospective cohort study from the 1st of January 2000 to the 15th of September 2016. We described epidemiology, clinical features, treatment and outcomes of patients admitted to ICU.

**Results** 19 patients were enrolled, the overall cumulative incidence was 0.2 episodes/1000 admissions and 0.17% of all community acquired pneumonia hospitalized during this period. Eighteen patients were males. Median age was 51 years (25–81 years). A summer and autumnal peaks was observed. Comorbidities were observed in 9 patients. Mean IGSII score was 42 (23–79) and major criteria of ATS were present in 5 cases. At admission in ICU, all patients had acute respiratory failure; 3 gastrointestinal symptoms; 9 hypotension, 8 rhabdomyolysis and 12 had elevated liver enzymes. Acute respiratory distress syndrome (ARDS) was present in 16 cases (severe \( n = 5 \), mild \( n = 6 \), light \( n = 5 \)). Septic shock and acute renal failure were observed respectively in 5 and 8 cases. Diagnosis was made in all cases by urinary antigen test. *Legionella pneumophila* was present also in sputum in 5 cases. Mechanical ventilation was needed in 13 cases; it was initially NIV in 46%. Fifteen patients were treated by a combination antibiotherapy: It was macrolide and quinolone in 52% (\( n = 10 \)), quinolone and rifampicine in 31% (\( n = 5 \)), the others were treated with macrolide or quinolones in monotherapy. Median duration of ventilation was 7 days (2–29 days). Median length of ICU stay was 13 days. Seven patients were died. Univariate analysis showed that presence of major criteria of ATS score at admission (\( p = 0.001 \)), severe ARDS (\( p = 0.004 \)), elevated liver enzymes (\( p = 0.047 \)), needs of mechanical ventilation (\( p = 0.024 \)) or intubation (\( p = 0.00 \)), septic shock (\( p = 0.001 \)); dialysis (\( p = 0.020 \)) were associated with a higher mortality. Multivariate analysis showed that independent predictive factors of mortality were ATS major criteria OR 3.5 [95% CI (1.08–11.92)]; mechanical ventilation OR 0.5 [95% CI (0.28–0.88)]; severe ARDS OR 0.117 [95% CI (0.017–0.80)]; septic shock OR 0.09 [95% CI (0.015–0.65)]; need of dialysis OR 0.14 [95% CI (0.02–1.06)].

**Conclusion** Our study confirms that LD requiring hospitalization in ICU is associated with high morbimortality. Presence of ATS major criteria, severe ARDS, mechanical ventilation, septic shock and dialysis were the major factors for worse prognosis.

**Competing interests**
None.

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**P132**

Acute respiratory failure in patients treated with chronic hemodialysis: pulmonary edema or not?

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Annals of Intensive Care 2017, 7(Suppl 1):P132

**Introduction** Patients on chronic hemodialysis are prone to develop overload pulmonary edema (OPE) but also infections, and have been reported to be regularly admitted to the intensive care unit (ICU) [1, 2]. Acute respiratory failure (ARF) in this specific population can be challenging for the intensivist, as OPE requires ultrafiltration which could be deleterious in other causes. The objectives of this study were (1) to describe the spectrum of ARF causes in patients on chronic hemodialysis and (2) to identify predictors of pulmonary edema diagnosis.

**Patients and methods** We performed a monocentric retrospective study in the 18-bed ICU of a non-teaching hospital with a nephrology unit. All patients on chronic hemodialysis admitted to the ICU for ARF

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Table 24 Characteristics at admission for SP patients and MP patients

| N (%) or median (IQR) | Mycoplasma pneumoniae patients (\( N = 76 \)) | Streptococcus pneumoniae patients (\( N = 76 \)) | \( \text{P value} \) |
|-----------------------|---------------------------------------------|---------------------------------------------|------------------|
| **Clinical respiratory findings** | | | |
| Respiratory rate | 33 [27–38] | 30 [26–36] | 0.43 |
| Signs of respiratory distress | 33 (49%) | 34 (45%) | 0.74 |
| Ronchi | 9 (15%) | 12 (16%) | 1 |
| Crackles | 36 (61%) | 44 (59%) | 1 |
| Signs of consolidation | 5 (9%) | 22 (30%) | 0.008 |
| Decreased vesicular sound | 10 (17%) | 28 (38%) | 0.007 |
| **Clinical presentation** | | | |
| Time since symptom onset (days) | 6 [4–9] | 3 [2–7] | 0.0008 |
| Fever | 58 (83%) | 54 (71%) | 0.12 |
| Shock | 6 (8%) | 24 (32%) | 0.0004 |
| Neurological disorders | 1 (1%) | 20 (26%) | <0.0001 |
| Gastro intestinal symptoms | 1 (1%) | 15 (20%) | 0.003 |
| **Radiological features** | | | |
| Number of quadrants | <2 | 37 (49%) | 66 (87%) |
| >2 | 16 (21%) | 9 (12%) |
| Alveolar opacities | 42 (75%) | 69 (92%) |
| Interstitial opacities | 20 (36%) | 6 (8%) |
| Pleural effusion | 3 (5%) | 17 (23%) |
in the period 2011–2015 were included. Patients with unknown diagnosis on ICU discharge were excluded. We defined ARF as either use of oxygen at ≥6 l/min dioxycyan to achieve a pulse oximetry measurement of ≥90% (88% in COPD patients) or partial pressure of arterial oxygen (PaO₂) <60 mmHg on room air or need for mechanical ventilation plus one of the following signs: a respiratory rate of >25 breaths/min, use of accessory respiratory muscles, or cyanosis associated to worsening dyspnea. The following data were collected: previous dry weight, time from last dialysis session, clinical, biological and imaging data on admission, number of dialysis sessions performed in ICU and total ultrafiltration volume required to withdraw oxygen therapy, weight on discharge, treatment and outcome. Patients were classified as having OPE (OPE group) or not (non-OPE group) based on the final diagnosis on ICU discharge. The two groups were compared with univariate analysis using non parametric tests.

Results 44 patients with 59 episodes of ARF were included in the final analysis. 46 (78%) episodes were due to OPE and 13 (22%) to other causes, including: 8 (61.5%) lower respiratory tract infections, 4 (30.8%) COPD exacerbation and one pulmonary embolism (7.2%). Patients admitted for OPE had higher systolic blood pressure at first medical contact (median [25–75 IQR]: 194 mmHg [176–215] vs 105 mmHg [91–119]; p < 0.0000001), lower CRP on arrival (10 mg/l [6–17] vs 53.5 mg/l [9.8–133.8]); p = 0.06), higher weight gain related to dry weight (2.1 [1–3] vs 0.5 [−0.25 to 2]; p = 0.01) and longer time since last dialysis (2 days [2–3] vs 1 [0–2.5]; p = 0.02). We did not find significant differences in temperature on arrival, white blood cells count nor NT-proBNP. Chest X-Ray interpretation was correct in 82% of cases of OPE patients but could not conclude in 46% of non-OPE episodes. Non-OPE patients had a more severe prognosis 38.5% of death (n = 5) versus 2.2% (n = 1).

Conclusion In this small retrospective monocentric study, ARF in patients on chronic hemodialysis is due to non-OPE cause in 22% of cases. A lower blood pressure, a higher CRP, and a shorter time from last dialysis session could prompt physician to search for a non-OPE cause and adjust cautiously ultrafiltration therapy.

Competing interests None.

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P133 "Urgent chemotherapy for life-threatening complications related to solid neoplasms"

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Introduction Both haematological and solid malignancies may be directly responsible for life-threatening organ failures including obstruction of anatomical structures, tissue infiltration, tumor lysis syndrome, or coagulation disorders. Besides advanced life support and eventual instrumental interventions, the treatment of cancer-related organ failures relies on timely administration of chemotherapy. Published data about requirements of chemotherapy in the ICU are mostly related to patients with haematological malignancies, while reports of patients with solid tumors are scarce. In this study, we addressed the features and outcomes of patients with organ failures directly related to solid neoplasms.

Patients and methods We performed a retrospective multicenter study within the GrrrOH research network. All adult patients who were admitted to the ICU with organ failures related to solid malignancies and treated with chemotherapy between 2000 and 2015 were enrolled into the study. Data were collected from individual files, and included the overall severity through the SOFA score computed at the time of ICU admission, the type and mechanism of organ failures, and the modalities of chemotherapy administration. Endpoints were the in-ICU and in-hospital vital status.

Results 136 patients were included. The most common underlying malignancy was lung cancer (n = 90), distributed between small cell lung cancer (SCLC) (n = 57) and non-small cell lung cancer (NSCLC) (n = 33). Most malignancies were newly diagnosed (n = 122, 89.7%), the diagnosis being made in the ICU for 80 patients (58.9%). The majority of patients (n = 82, 60.3%) had metastasis. The main reason for ICU admission was acute respiratory failure in 111 (81.6%) patients. Compression and tissue infiltration by tumor cells were the leading mechanisms resulting in organ involvement in 78 (57.4%) and 47 (34.6%) patients. Other indications for ICU admission relied on paraneoplastic manifestations in 11 cases. The treatment was based on combined and single chemotherapy in 120 (88.2%) and 16 (11.8%) patients, respectively. The dosing was reduced in 20 patients (20%) mostly linked to renal failure, and the sequence of drug administration was modified in 18 patients (18%). Thirty-four patients (25%) required either instrumental or surgical adjuvant procedures. Eighty-nine patients received invasive mechanical ventilation with duration of 13 (5–25) days, 39 patients required vasopressors (28.7%) and 11 required renal replacement therapy (8.1%).

The overall in-ICU, in-hospital, 6-month and 1-year mortality rates were 37, 58, 74 and 88%, respectively. In a multivariate logistic regression analysis, SCLC was identified as an independent predictor of hospital survival. However this gain in survival was not sustained since the 1-year survival rates of SCLC, NSCLC and non-lung cancer patients all dropped below 20%.

Discussion The prognosis of solid neoplasm-related organ failures relies on multidisciplinary management involving both intensivists and oncologists. Appropriate management of cytostatic drugs is paramount to improve their efficacy on tumor cells while minimizing their toxicity. Dosing of chemotherapy in critically ill patients represents a still unexplored area of research owing to a number of factors that may result in drug underdosing or overdosing.

Conclusion Acute respiratory failure related to lung cancer represents the main indication for urgent administration of chemotherapy in the ICU. Urgent chemotherapy along with aggressive management of organ failures in the ICU can be life-saving in a number of cancer patients, most especially for SCLC, although the long-term survival is hardly sustainable.

Competing interests None.
Lung ultrasound for early diagnosis of pneumonia after cardiac surgery

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Introduction Pneumonia is a frequent and severe complication of major cardiac surgery, contributing to postoperative morbidity and death [1]. Diagnosis of pneumonia remains a challenge in ventilated patients, notably after cardiac surgery, and the diagnostic performance of Clinical Pulmonary Infection Score (CPIS) remains controversial. Lung ultrasound (LUS) has been successfully used for the diagnosis and the management of community acquired pneumonia and ventilator-associated pneumonia (VAP) [2], but LUS usefulness and reliability was never investigated in the specific ICU patients after cardiac surgery with cardiopulmonary bypass (CPB).

This pilot observational study investigated the clinical relevance of lung ultrasonography (LUS) for diagnosis of pneumonia in cardiac postoperative patients with acute respiratory failure (ARF).

Patients and methods Adult patients were prospectively enrolled from January through May 2015 in presence of acute respiratory failure (ARF) less than 3 days after a cardiac surgery with CPB. Lung ultrasound examination was performed as follows: subpleural consolidation, lobar consolidation, dynamic and static air bronchogram, and interface of pleural effusion. A new score, combination of simplified CPIS and a simplified CPIS was used for the diagnosis of pneumonia in ARF patients with an inexpensive, non-invasive and convenient at bedside technology.

Conclusion This prospective observational study is the first one showing that LUS combined with a clinical score can be a reliable tool for early diagnosis of pneumonia in a cardiac ICU population after cardiac surgery with CPB.

Competing interests None.

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Trends in intensive care admission for respiratory infections attributable to the elderly in aging population

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Introduction Acute respiratory infection (ARI) is the most common infectious cause for admission to Intensive Care Unit (ICU) and the prevalence of infection increases with age. Changes in population demographics and comorbid illness may drive important changes in the composition of patients admitted to the ICU. Notably, how the population aging impacts on the incidence of hospitalized patients for ARI and how it increases the demand for critical care services is unknown. To address this knowledge gap, we sought to describe trends in demographics changes among elderly patients admitted to ICU for ARI on a 9-year period.

Materials and methods We conducted a retrospective cross-sectional study based on hospital discharge databases (HDDs) from January 1, 2006 to December 31, 2014. We selected patients over 18 years old (y-o) who were hospitalized for ARI in a French region (Centre Val de Loire region, 2.5 million inhabitants, served by one university hospital, one regional hospital and 37 general and private hospitals). Cases of ARI were extracted from the HDD with an algorithm based on ICD-10 specific diagnosis codes, taking into account the type, number and position of these codes in the hospital discharge report. We previously validated the ICD-10 case definition reviewing a sample of medical charts as the gold standard. Giving the acceptable accuracy and precision of our algorithm, the following data were extracted from the
Pulmonary embolism in intensive care unit: incidence and impact prognosis

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Introduction

Pulmonary embolism is a major complication observed in critically ill patients. ICU environment patients are at high risk of deep vein thrombosis and pulmonary embolism. The objective of our study is to determine the incidence of this pathology and to analyze the prognosis of patients having a pulmonary embolism in ICU.

Patients and methods

Prospective study included all patients with confirmed pulmonary embolism by a spiral computed tomography scan showing one or more filling defects in the pulmonary artery or in its branches, over a period of 1 year from 1 July 2014 to 30 June 2015. We studied the epidemiological, clinical, biological and radiological features of these patients. We compare two groups in univariate analysis: survivors and deaths.

Results

During the study period, 705 patients were admitted in our ICU. The diagnosis of PE was confirmed in 75 patients (10.6%). The mean age was 63 ± 15 years. The sex ratio (M/W) was 75% and 25%, respectively. The mean SAPS II score on ICU admission was 37 ± 16. The mean length of stay was 10 ± 7 days. The mean PaO2/FiO2 was 200 ± 50 mmHg and the presence of shock. The mortality rate in ICU was 25%.

Conclusion

Pulmonary embolism (PE) is a frequent and aggressive complication in critically ill patients. ICU environment patients are at high risk of deep vein thrombosis and pulmonary embolism. The objective of our study is to determine the incidence of this pathology and to analyze the prognosis of patients having a pulmonary embolism in ICU.

We found an increasing incidence of PE over the study period: 5% in 2006 whereas it was superior to 13% since 2010. This trend for increase ICU admission was proportional with aging: 2-fold increase at 75–79 y-o, 2.5-fold increase at 80–84 y-o, threefold increase at 85–90 y-o, and 4-fold increase for patients over 90 y-o. The overall hospital-mortality was relatively stable and varied between 6.7% and 9% for non-ICU patients and 14.4–17.2% in ICU patients. Regarding patients over 90 y-o hospitalized in ICU, the mortality has been drastically reduced: from 40.9% to 22.2% on the study period.

Conclusion

This large prospective study provides a comprehensive view of elderly with severe ARI admitted to ICUs. We found significant increase in elderly patients with primary diagnosis of ARI hospitalized in ICU. We highlighted that ARI is an increasing reason for death among ICU-hospitalized patient and still a major public health problem. Interestingly, our results are consistent with longitudinal changes in ICU admission also observed in this study period in the United States.

Conclusion

We found substantial increase of ARI diagnoses leading to hospitalization between 2006 and 2014 with a growing demand for critical care service. Further studies are needed to evaluate the benefits of intensive care hospitalization for the very elderly patients. We cannot arbitrary reject the very elderly from the doors of the ICU—it is ageism—but we need information to select more accurately the patients with the highest probability of survival, and to avoid useless and aggressive cares when it is not appropriate.

Competing interests

None.

P138

Correlation between the transcranial color-coded duplex sonography and the optic nerve sheath diameter in the prediction of intracranial hypertension

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Introduction

Intracranial hypertension is suspected on clinical, radiological and ophthalmological criteria. The measurement of intracranial pressure is the reference method. Due to infectious and hemorrhagic risks, other non-invasive techniques have been developed such as the transcranial color-coded duplex sonography (TCCDS) and the optic nerve sheath diameter (ONSD). The aim of our study was to compare TCCDS data to the ONSD measurements and to CT scanning data in predicting intracranial hypertension.

Patients and methods

Prospective study enrolling all patients admitted to our ICU for severe traumatic brain injury between February and August 2016. The intracranial hypertension was studied by CT scanning, TCCDS and ocular ultrasonography. CT defines intracranial hypertension by a midline shift >5 mm, collapsed third ventricle, presence of hydrocephalus and erased basal cisterns with significant edema. TCCDS identifies a significant intracranial hypertension by a pulsatile index (PI) >1.2 and a diastolic velocity <20 cm/s. An ONSD > 5 mm is considered abnormal. The study of associations was done using non-parametric tests (Wilcoxon and Fisher) and data analysis was performed by epi-info software.

Results

Sixty patients with severe traumatic brain injury were included in the study. The average age was 42 ± 17 years. The sex-ratio was 1.4. The intracranial hypertension was diagnosed by pectillar abnormalities in 36% of cases, CT scanning in 84% of cases, TCCDS in 68% of cases and ONSD in 69% of cases. There was no significant correlation between an ONSD > 5 mm and CT scanning findings (p = 0.58) nor with TCCDS (p = 0.08) in predicting intracranial hypertension.

Conclusion

ONSD is a quick and simple method that has been widely used for predicting intracranial hypertension. A larger sample is needed to confirm our results.

Competing interests

None.

P140

Traumatic brain injury caused by traffic accidents: epidemiology and prognostic factors: a multivariate analysis of 694 cases

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Introduction

Traffic accidents are responsible for 1.4% of outpatient consultations and constitute 24% of hospitalisations in ICU. Moreover, comparison between survivors and deceased showed that factors associated with deaths were: high SAPS II score on ICU admission, high SAPS III score on ICU admission, high SOFA score, low shock index on the day of pulmonary embolism, hypoxemia with PaO2/FiO2 < 200 mmHg and the presence of shock. Prevention is advised.

Competing interests

None.
Introduction Head injuries are a public health problem. Studies on the epidemiology and especially the prognosis of head trauma are rare in developing countries. The objective of our study is to determine the epidemiological aspects and to identify factors correlated with short- and long-term prognosis.

Patients and methods It was a retrospective study during 4 years (January 2009–December 2012) included 694 patients with traumatic brain injury due to traffic accidents. We studied the factors correlated with poor prognosis in terms of death or Glasgow Outcome Scale (GOS) in univariate and multivariate analysis.

For study according to GOS, patients (n = 694) were divided in two groups: favorable GOS for patients having a good recovery or recovery with a minor handicap [GOS class 4 or 5] and unfavorable GOS for patients having a severe handicap or vegetative state [GOS class 1 or 2] or died (GOS1).

Results During the study period, 694 patients (18.7% of all patients) were admitted to our ICU with traumatic brain injury due to traffic accidents. The mean of age was 31.9. The sex ratio was 5.8. In admission, 20% of patients had hypotension, 17.4% of patients had respiratory distress and the mean GSC was 8.8. The mortality rate was 28.5%. 496 patients were survivor: 13 patients with vegetative state, 133 with severe handicap (19.3%), 185 patients with minor handicap (26.7%), and 163 patients with good recovery (23.6%). The predictive independent factors correlated to mortality in multivariate analysis were: age &gt;= 38, hypotension, hypoxemia, SGS &lt; 8, unilateral or bilateral mydriasis, cerebral edema, Marshall Class VI, initial hemoglobin &lt; 11.3, Blood sugar &gt;= 8.3 mmol/l. The predictive factors correlated with poor prognosis according GOS were: age &gt;= 38, initial shock, SGS &lt; 8, unilateral or bilateral mydriasis, post traumatic coma duration more than 5 days, cerebral edema, initial hemoglobin &lt; 11.7 g/dl, blood sugar &gt; 8.4 mmol/l and catecholamine using.

Conclusion The short-term prognosis of head trauma seems recently stabilized even for the most serious patients, but the distant consequences of head trauma are fairly frequent, heavy and often undervalued. That is why, it is so important to identify these consequences and management them correctly.

Competing interests None.

P141 Benefice of rib fixation in chest trauma: a before/after study Pierre-Julien Cungi1, Cédric Nguyen1, Jean Cotte1, Erwan D’aranda1, Eric Meaudre2, Jean-Philippe Avaro2
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Introduction Fifteen percent of the trauma has a chest trauma. Depending on the type of injury, mortality raises from 4 to 60%. There are no surgical guidelines concerning rib fracture except for flail chest needing mechanical ventilation. No trial has yet assessed the global benefit of both surgical and medical treatment. The primary endpoint was to determine wether rib fixation on the one hand and epidural analgesia on the other hand reduces pneumonia. The secondary endpoints were to show a decrease of mechanical ventilation duration and lengths of stay in intensive care and hospital.

Materials and methods We performed a pre/post osteosynthesis study. We determined two groups. From 01/2011 to 08/2014, all patients admitted for chest trauma were screened by a thoracic surgeon in order to select those with a rib fixation indication. They formed the historical cohort: «not fixed». From 08/2014 to 08/2016, all patients with chest trauma and rib fixation were included. They formed the group «fixed». Were excluded patients who died within the first 48 h. The surgical treatment is defined by a rib fixation. The medical treatment is defined by epidural analgesia and noninvasive ventilation.

Results One hundred and twelve patients with equivalent surgical indication of rib fixation were enrolled from January 2011 to August 2016. Fifty-seven patients were not fixed and 55 patients were fixed. Eighty-two (73%) were men of 57 (20–86) years old. The mean IGS was 22 (4–66). The mean SAPS II was 25 (6–86). Eighty-five patients (80%) were hospitalized in Intensive care. 72 patients (64%) had an epidural analgesia. Fifty percent of the patients were operated on within the first 24 h. Thirty-three patients suffered from pneumonia that occurred on average on the fifth (5–10) day. Epidural analgesia significantly decreased the incidence of pneumonia (p &lt; 0.001). Rib fixation had no impact on the incidence of pneumonia (p = 0.1). Both length of stay in hospital and in intensive care were neither reduced by epidural analgesia nor by rib fixation. Epidural analgesia reduced mechanical ventilation duration. Rib fixation didn’t reduce mechanical ventilation duration.

Conclusion It is the first study to assess the importance of both medical and surgical care of chest trauma. Epidural analgesia is effective in decreasing incidence of pneumonia whatever the rib fixation.

Competing interests None.

P142 Prognostic factors of severe liver injury Med Azz Bouhouri1, Mohamed Taouftik Slaoi1, A. Soufi1, K. Khaleq1, D. Hamoudi1, A. Nsiri1, R. Harrass1
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Introduction Serious liver injuries classified have very high mortality figures despite improved care systems and means of paraclinical investigations.

Stud Objective: The purpose of this study was to determine the prognostic factors of severe liver injury by evaluating epidemiological, diagnostic, therapeutic and evolutionary data and this to promote conservative therapeutic management.

Patients and methods In this retrospective clinical study over a period of 3 years (May 2013–May 2016), patients admitted to the intensive care surgical emergencies Ibn Rochd CHU Casablanca for serious liver injury were evaluated on specific criteria for inclusion and exclusion in using a pre-established operating profile including epidemiological, clinical, biological, radiological and therapeutic findings with a significance level set at 5%.

Results After this study, 23 patients were compiled. 19 patients were male and 4 female, mean ages was 34 ± 11.7 years with a range from 17 to 58, the etiologies were dominated by accidents of the high way followed by assaults and finally fall from heights. An unstable hemodynamic status was initially found in 6 cases (26%) and the use of vasoconstrictor was necessary in 18 patients (78.2%). Ultra sonography was performed in 16 patients (69.5%) and objectified isolated hem peritoneum or associated with visceral lesions in all patients. Abdominal CT scan was performed to 20 patients (87%) objectifying in 10 cases (50%) different findings with those of ultrasound.

An emergency surgery was raised in 15 cases and delayed surgery was indicated in 3 patients. Mortality in our study was 39.13% where hemorrhagic shock presents the leading cause. The prognosis was closely linked to the value of systolic blood pressure at admission, the SPO2, the Glasgow coma scale, haemoglobin value, the presence of hem peritoneum of great abundance and whether or not a use of vasoconstrictor.

Conclusion The development of imaging techniques in emergency, especially FAST ultrasound and computerized tomography scan, have modified diagnostic and therapeutic attitude in severe liver injury over the last years. The yellow a conservative approach based on the stability of the hemodynamic status and the response to new resuscitation techniques, this approach requires strict supervision and...
involves a multidisciplinary team (intensivist, surgeon and radiologist) available in emergency for cases with complications.

Competing interests
None.

P143
Acute burn care: A mythe or a reality?
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Introduction
Early burn resuscitation of major burn-injured patients is the cornerstone of burn care and aims to improve outcome and decreases morbidity and mortality rates of these patients. So, initial care of severely burned patients requires participation of all physicians in every discipline and in every hospital. The goal of this study was to examine characteristics of burn patients acutely transferred to our intensive burn care unit and to assess their prognosis.

Patients and methods
A prospective study was conducted in intensive burn care center in Tunis. All consecutive adult burned patients acutely transferred to our burn center, from January 1st to September 23rd, 2016 were included in the study. Demographic, clinical and biological data of patients were recorded.

Results
During the 9 month study period, 190 patients were admitted among which 101 patients were acutely transferred from other hospitals (53%). The mean age was 37 ± 15 years. The mean surface burned area announced was 44 ± 22 versus 36 ± 22% reevaluated at admission. Patients were transferred with a delay of 40 H after burns [H1–H264]. Burn injuries were caused by domestic accidents in 42%, self immolation in 26% and work related burns in 14%. Transfer with medical agreement was noted in 57% of cases. At admission, 12% of patients had burn shock and 43% had endotracheal intubation. A central venous catheter was placed in 51% of cases, nasogastric tubes in 11% and urinary devices in 55% of cases. Dressing were performed in 66% of cases. Fluid resuscitation was initiated in 74% of cases with crystalloid: Ringer lactate (42%) and/or normal saline (19%). Initial lactate of patients was 3.36 ± 1.7 mmol [1–8.8] with pH at 7.33 [6.6–7.51] and bicarbonates at 20 ± 5 [9–34]. We noted that patients transferred without medical agreement had more burn shock (16.2 vs 8.6%) and a higher mortality (25.5 vs 17.2%).

Conclusion
Early critical care of severely burned patients, especially, fluid resuscitation and monitoring, coupled with appropriate early referral to a specialist, greatly help in minimizing complications and optimizing prognosis.

Competing interests
None.