Hyperlactatemia in patients with non-acetaminophen-related acute liver failure

Pilar Taurá, Graciela Martinez-Palli, Julia Martinez-Ocon, Joan Beltran, Gerard Sanchez-Etayo, Jaume Balust, Teresa Anglada, Antoni Mas, Juan-Carlos Garcia-Valdecasas

Abstract

AIM: To characterize hyperlactatemia in patients with non-acetaminophen acute liver failure (ALF) in an attempt to clarify the mechanisms implicated and the role as a prognosis factor.

METHODS: In the setting of liver transplantation, 63 consecutive patients with non-acetaminophen acute liver failure were studied in relation to tissue oxygenation, hemodynamic and metabolic parameters. Before and after transplantation, the number of infected patients and outcome were registered.

RESULTS: Acute ALF showed higher levels of lactate than subacute ALF (5.4 ± 1 mmol/L versus 2.2 ± 0.6 mmol/L, P=0.01). Oxygenation parameters were within the normal range. Lactate levels showed good correlation with respiratory quotient (r=0.759, P<0.005), mean glucose administration (r=0.664, P=0.01) and encephalopathy (r=0.698, P=0.02), but not with splanchnic arteriovenous difference in PCO2, pH and the presence of infection (P=0.1). Portal vein lactate was higher (P<0.05) than arterial and mixed venous lactate, suggesting its production of hyperlactatemia in the intestine and spleen. The presence of infection was an independent predictor of survival.

CONCLUSION: Hyperlactatemia is not a prognosis factor due to byproduct of the overall acceleration in glycolysis.

Key words: Hyperlactatemia; Non-acetaminophen acute liver failure; Splanchnic hypoperfusion; Acute liver failure

Taurá P, Martinez-Palli G, Martinez-Ocon J, Beltran J, Sanchez-Etayo G, Balust J, Anglada T, Mas A, Garcia-Valdecasas JC. Hyperlactatemia in patients with non-acetaminophen-related acute liver failure. World J Gastroenterol 2006; 12(12): 1949-1953

http://www.wjgnet.com/1007-9327/12/1949.asp
lactate levels as well as the role as a prognosis factor.

MATERIALS AND METHODS

Patients
A total of 63 patients with diagnosis of non-acetaminophen ALF were admitted to the Liver Intensive Therapy Unit in our hospital and submitted to liver transplantation. All patients were prospectively evaluated and included in the study. Four of them were considered hyperacute, 39 acute while 20 subacute hepatic failure based on the criteria of the Kings College Hospital Group[8].

This prospective study was approved by the Clinic Hospital Research Ethic Committee. Informed consent for the study was obtained from each patient or patient’s family.

General management
All patients were managed in the Liver Intensive Therapy Unit, in a standard manner[7]. They were maintained on 10%-20% glucose solution infusion (keeping blood glucose levels between 90 and 120 mg/dl). Prophylaxis for upper gastrointestinal bleeding and close microbiological surveillance (all patients received norfloxacin and nystatin) were performed. Coagulation profile was corrected only in the presence of overt bleeding. Those in grade III and IV coma (43 patients) were treated with standard sedation and mechanical ventilation. An epidural transducer (Ladd Research Laboratories, Burlington, VT) was inserted into the epidural space to enable early recognition and treatment of intracranial hypertension (>25 mmHg), mannitol (0.5 to 1 mg/kg) for over 30 min and pentobarbital infusion as a following step were used to control intracranial pressure (ICP). Systemic arterial hypotension (systolic <100 mmHg) was managed by dopamine infusion according to the hemodynamic conditions in an attempt to maintain cerebral perfusion pressure (CPP) (>60 mmHg). No artificial liver support techniques were used and no patient was treated with N-acetylcysteine or epoprostenol.

Anesthesia and hemodynamic management
In 20 patients who remained awake (encephalopathy grades I and II), anesthesia was induced following our standard policy. All patients were mechanically ventilated (Servo 900C, Siemens) and the fraction of inspired oxygen was adjusted to achieve an arterial PaO₂ of 180 - 220 mmHg. Ventilatory parameters were regulated to maintain the end tidal CO₂ around 32 mmHg and P:CO₂ below 35 mmHg. In all patients vasopressor dopamine at 3 µg/kg was administered from the beginning of the surgical procedure and maintained through the transplant and increased to 8 µg/kg if necessary to achieve a mean arterial pressure greater than 75 mmHg. After graft reperfusion, arterial hypotension was treated by intravenous bolus of 10 µg of epinephrine. Fiberoptic pulmonary artery floatation catheter (7.5 French, Edwards Laboratories, Irvine, Calif.) was introduced through the right internal jugular vein and an arterial catheter (18 gauge, Arrow, Reading, PA) was placed via the left radial artery. At the beginning of the surgical procedure portal vein catheter (Certofix® Mono S330 16G Braun) was placed by introducing it through a branch of the superior mesenteric vein and advanced to the portal vein until it was felt, in order to register portal pressure and to obtain blood samples for oxygen and metabolic parameters in splanchnic area. Not all patients received portal vein cannulation because it was not considered technically possible by the surgeon in 11 patients (4 with subacute ALF and 7 with acute ALF).

Variables analyzed in the setting of LTx
Before the surgical transplant procedure was started, the following parameters were measured. Mean arterial pressure (MAP, mmHg), cardiac index (CI, L/min/m²) and systemic vascular resistance index (SVRI, dyn.sec.cm⁻³.m⁻¹) were obtained.

Systemic parameters including oxygen content difference [D(a-v)O₂, ml/dl], oxygen delivery (DO₂, ml/L/min/m²), oxygen consumption (VO₂, ml/L/min/m²), oxygen extraction ratio (VO₂/DO₂, %), mixed venous/arterial gradient of PCO₂ (VACO₂, mmHg), arterial/mixed venous gradient of pH (AvPHe, U) as well as respiratory quotient (RQ) were calculated.

Splanchnic parameters including arterial/portal venous oxygen content difference [D(a-v)O₂, ml/dl], oxygen extraction ratio (VO₂/DO₂), portal veno/arterial gradient of PCO₂ (VACO₂, mmHg), and arterial/portal venous gradient of pH (AvPHe, U) were obtained. All these variables were calculated following standard formulas.

Plasma lactate levels were measured with three blood samples simultaneously drawn from arterial catheter (IA), the distal part of pulmonary catheter (Lv) and portal vein catheter (Lp). Blood lactate level was determined using a Kodak Ektachem 700XR (Rochester, NY, USA) analyzer[8].

The need of catecholamine administration and the mean glucose administration in the last 48 hours before transplant were recorded. The need of vasopressor drugs administered during the transplant procedure was also registered. The degree of encephalopathy and intracranial pressure were recorded.

Explanted liver pathology
Weight and structural characteristics of all explanted livers were studied.

Infectious complications
Diagnosis of infection was made by the presence of the white blood cell count greater than 12×10⁹/L or less than 4×10⁹/L, the presence of immature neutrophils, temperature higher than 38 °C or lower than 36 °C and microbiological confirmation. Also, chest infection was confirmed by radiology. All these parameters were recorded daily during the ICU admission before transplantation and the ten following days. The number of infected patients and episodes and pathogens microbiologically confirmed was registered, respectively.

ICU stay and immediate outcome
The mean stay in ICU during the first admission of patients after transplantation and immediate outcome (first admission in the hospital) were recorded.
**RESULTS**

Demographic and clinical characteristics of the patients are shown in Table 1. No patient needed blood transfusion before the transplant.

**Systemic hemodynamic and oxygenation parameters**

Systemic hemodynamic parameters showed a hyperdynamic circulatory status with high CI (4.53 ± 1.4 L/min/m²) and low SVRI (1029 ± 420 dyn.sec.cm⁻⁵.m²). Oxygenation parameters remained within normal ranges (DO₂: 623 ± 36 mL/min/m², VO₂: 96.4 ± 21 mL/min/m² and VO₂/DO₂: 18.4 ± 3.1%).

**Blood lactate levels**

The production of lactate in the intestine or in the spleen (Table 2), in accordance with Murphy et al.⁶, was suggested by the increased difference between portal, arterial and mixed venous lactate (P < 0.05). Acute ALF showed significant higher levels (acute: 5.4 ± 1 mmol/L, subacute: 2.2 ± 0.6 mmol/L, P = 0.01).

Lactate levels did not correlate with any of the hemodynamic or oxygenation parameters studied, except for the respiratory quotient (Figure 1 A). The grade of encephalopathy and the amount of glucose administered 48 hours before transplant (Figure 1 B and Figure 1 C) correlated well with blood lactate levels.

**Statistical analysis**

Statistical analysis was performed using two-sided paired t-test for comparison of paired data and two-sided unpaired t-test for comparison of groups. The Bonferroni correction test was applied as appropriate. Categorical data were compared with the chi-square test for relationship between encephalopathy and infection. Correlation between lactate and the hemodynamic and oxygenation variables was assessed by linear regression analysis. Variables reaching significance in the univariate analysis between survivors and non-survivors were included in the multivariate analysis. Multivariate analysis was carried out by stepwise logistic regression analysis to determine discriminants of survival. All values were expressed as mean±SD. P < 0.05 was considered statistically significant.

**Table 1 Clinical and demographic characteristics of the patients**

| Age (yr) | 32.7 ±11 |
| Sex (M/F) | 26/37 |
| Etiology (n, %) |  
| Viral hepatitis | 18 (30) |
| Cryptogenic (non-A, non-B, non-C) | 32 (52) |
| Drug toxicity | 7 (11) |
| MAOI | 5 |
| Rifampin+Isoniacid | 2 |
| Isoturitarine | 1 |
| α methil-dopa | 1 |
| Metabolic disease | 4 (6) |
| Encephalopathy |  
| I-II (subacute form: 20 p) | 20 (31.7) |
| III-IV (acute form:39 p) | 43 (68) |

**Table 2 Plasma lactate levels at different sites (mean±SD)**

| Variable | Total (n=52) | Acute (n=36) | Subacute (n=16) | P |
|-----------------|--------------|--------------|-----------------|---|
| La (mmol/L) | 4.1 ±1.8 (1.1-15.2) | 5.4 ±1 | 2.2 ±0.6 | 0.01 |
| Lv (mmol/L) | 4.3 ±2 (2.4-16.3) | 4.7 ±2 | 2.4 ±1.4 | 0.03 |
| Lp (mmol/L) | 5.3 ±1.1a (2.1-17.6) | 6.8 ±1.8a | 2.8 ±0.9 | <0.01 |
| pH (units) | 7.36 ±0.07 (7.28-7.42) | 7.33 ±0.02 | 7.36 ±0.04 | NS |
| VACO₂(mmHg) | 12.4±7 (8.3-13.4) | 13.8±5 | 9.6±9 | NS |
| AvpH (units) | 0.06 ±0.03 (0.02-0.07) | 0.08±0.02 | 0.07±0.04 | NS |

**Splanchnic oxygenation parameters**

The splanchnic VACO₂ and AVpH levels were in normal range and did not correlate with plasma lactate levels (r = 0.203 and r = 0.164, respectively). No differences were found (Table 2) when we compared patients with high (acute ALF) and low lactate levels (subacute ALF).

In order to maintain hemodynamic stability, only five patients with acute ALF needed administration of dopamine prior to the transplant. Only one of the patients demonstrated a high level of lactate (8.2 mmol/L). There was no significant correlation between lactate levels and the dosages of dopamine and epinephrine (r = 0.020 and r = 0.13 respectively) through the transplant. The total dose of epinephrine administered after graft reperfusion in patients with low (subacute ALF) and high lactate level (acute ALF) was similar in both groups (42.6 ± 12 µg and 54.1 ± 18 µg, respectively).

**Explanted liver pathology**

No relationship was found between lactate levels and liver weight. Interestingly, although the rate of massive necrosis was similar, liver weight of subacute ALF patients was significantly lower than that of acute ALF patients (762 ± 22 g and 932 ± 38 g, P < 0.05).

**Infectious complications**

Twenty-one patients (33.3%) were infected in the perioperative period. Bacterial infection was found in 16 patients (25.4%), fungal infection in 5 patients (7.9%) and viral infection in 2 patients (3.1%). The incidence of infectious episodes of acute (13 patients, 30.2%) and subacute (8 patients, 40%) hepatic failure showed no difference. The level of lactic acidosis (Figure 2) no correlated with the presence of infection (P = 0.1). Sixteen patients were infected before the transplant, 11 of them requiring mechanical ventilation because of pulmonary infection (confirmed on chest radiograph and microbiologically), showed significantly higher arterial lactate levels compared with mixed venous lactate level (5.2 ± 1.1 and 4.5 ± 0.8 mmol/L, P = 0.03), suggesting lactate production within the lungs.

**ICU stay and immediate outcome of patients with ALF**

During the stay in ICU, 13 patients died (8 patients with acute and 5 subacute liver failure). Bleeding was not controlled in 2 patients. Furthermore, 3 patients needed
Arterial lactate levels in 21 infected and 40 non-infected patients. Box n
Correlation between arterial lactate level (nmol/L) and respiratory quotient (www.wjgnet.com
It is important to emphasize that the foremost cause with the degree of hyperlactatemia.

DISCUSSION

Hyperlactatemia observed in the context of non-acetaminophen ALF seemed to be related to aerobic glycolysis but not to tissue hypoperfusion. Additionally, the incidence of infection in these patients, which appeared as an independent predictor of survival, did not correlate with the degree of hyperlactatemia.

It is important to emphasize that the foremost cause of ALF in our series was acute viral hepatitis that differs substantially in other series. Since acetaminophen exerts a direct toxic effect on cellular respiration, this type of cytotoxicity may be different from other types of ALF.

Currently, the presence of hyperlactatemia in patients with ALF is assumed as the consequence of underlying overt tissue hypoxia. In these patients, oxygen delivery may be impaired. Combined measurement of VACO2 and AVpH may serve as a good indicator of tissue hypoxia with a closer relation to cardiac output than blood lactate, because it is less affected by changes in fuel substrate utilization or enzymatic alterations [10-12]. In our study no differences were found in splanchnic VACO2 and AVpH between patients with high (acute ALF) and low lactate levels (subacute ALF).

If hyperlactatemia is attributable to tissue hypoperfusion, the increase in oxygen delivery should reduce lactate levels. However, several reports [13-15] have failed to identify this evidence in patients with ALF due to acetaminophen overdose.

In our patients selected to receive liver transplantation, plasma lactate levels did not correlate with hemodynamic or oxygenation parameters, except for RQ and the amount of glucose administered, suggesting that aerobic glycolysis is responsible of lactate hyperproduction. Since patients with ALF may exhibit hyperinsulinemia due to pancreatic hypersecretion and/or decreased hepatic clearance of insulin, they need glucose infusion in order to maintain blood glucose level. Previous studies [16-18] showed that infusion of glucose results in a dose-dependent rise in splanchnic lactate levels. Moreover, peripheral tissues of cirrhotic patients produce an exaggerated lactate production in response to glucose administration [19]. In our study, 7 patients who did not require glucose administration (subacute liver failure), showed normal lactate levels (<1.3 mmol/L). A complete round of the Cori cycle is proton-neutral because the H+ produced by lactate from glucose is subsequently removed during synthesis of glucose from lactate in the liver. In the context of liver failure, the possibility to handle lactate to glucose (gluconeogenesis) is handicapped and consequently the possibility to develop “lactic acidosis” without the presence of high muscular lactate overproduction is not significant. In our patients the arterial blood pH stayed within normal ranges (7.28 to 7.41). If hyperlactatemia is not related

www.wjgnet.com
to the mortality, infection, or operative hemodynamic management difficulty, the usage of dichloroacetate (which acts by stimulating pyruvate dehydrogenase) is not a priority unless its benefit to avoid postoperative severe alkalosis (lactate metabolism by the graft consume $H^+$) is considered. However, it seems that this treatment fails to attenuate metabolic alkalosis after transplant$^{[20]}$.

Clemmesen et al$^{[6]}$ have suggested a hypermetabolic condition secondary to an excessively high glycolysis in relation to the small liver mass, but we have not shown any correlation between liver weight and splanchnic lactate level in patients with ALF. Wendon et al$^{[21]}$ demonstrated that there is a significant correlation between arterial lactate and survival in acetaminophen related ALF, but in our study no relationship was found between plasma lactate and outcome. Inadequate splanchnic blood flow and tissue hypoperfusion could contribute to bacterial translocation and sepsis, however in these patients several factors can predispose to infection such as comatose state, steroid therapy, mechanical ventilation and intravenous catheters. Patients with ALF are susceptible to infection as a result of multiple immunologic defects (excessive cytokine production from cells, such as monocytes and macrophages in response to a number of stimuli including bacterial lipopolysaccharide)$.^{[22]}$. In our patients the Gram-positive organisms were predominant. The presence of Gram-negative organisms such as pseudomonas was not uncommon, showing a high incidence in this group of patients. These results suggest that selective bowel decontamination can be used in the treatment of such patients$.^{[23]}$. We did not find any correlation between lactate levels and infection incidence although there is evidence that cytokines may promote augmented production of lactate$^{[24]}$ by several organs, enhancing cellular glucose uptake and glycolytic metabolism. However, in agreement with other studies$^{[25]}$, arterial lactate is higher than venous mixed lactate in patients with pulmonary infection. In summary, the presence of high plasma lactate levels in patients with non-acetaminophen ALF, is not related to tissue hypoperfusion and is not a prognostic factor in the treatment of ALF.

REFERENCES

1. Makin AJ, Hughes RD, Williams R. Systemic and hepatic hemodynamic changes in acute liver injury. Am J Physiol 1997; 272: G617-G625
2. Bihari D, Gimson AE, Lindridge J, Williams R. Lactic acidosis in fulminant hepatic failure. Some aspects of pathogenesis and prognosis. J Hepatol 1985; 1: 405-416
3. Bihari D, Gimson AE, Waterson M, Williams R. Tissue hypoxia during fulminant hepatic failure. Crit Care Med 1985; 13: 1034-1039
4. Clemmesen JO, Gerbes AL, Gürberg V, Hansen BA, Larsen FS, Skak C, Tygstrup N, Ott P. Hepatic blood flow and splanchnic oxygen consumption in patients with liver failure. Effect of high-volume plasmapheresis. Hepatology 1999; 29: 347-355
5. Clemmesen JO, Høy CE, Kondrup J, Ott P. Splanchnic metabolism of fuel substrates in acute liver failure. J Hepatol 2000; 33: 941-948
6. O’Grady JG, Schalm SW, Williams R. Acute liver failure: redefining the syndromes. Lancet 1993; 342: 273-275
7. Castells A, Salmerón JM, Navasa M, Rimola A, Solé J, Andreu H, Mas A, Rodés J. Liver transplantation for acute liver failure: analysis of applicability. Gastroenterology 1993; 105: 532-538
8. Krofp J, Marx AM, Hildebrandt J, Gressner AM. Practical implications of coexistent different technologies in clinical chemical laboratories. Solid phase chemistry and conventional analysis. Eur J Clin Chem Clin Biochem 1991; 29: 675-683
9. Murphy ND, Kodakat SK, Wendon JA, Jooste CA, Muiseman P, Rela M, Heaton ND. Liver and intestinal lactate metabolism in patients with acute hepatic failure undergoing liver transplantation. Crit Care Med 2001; 29: 2111-2118
10. Ducy JP, Lamiiel MJ, Gueller GE. Arterial-venous carbon dioxide tension difference during severe hemorrhage and resuscitation. Crit Care Med 1992; 20: 518-522
11. Bakker J, Vincent JL, Gris P, Leon M, Coffenils M, Kahn RJ. Veno-arterial carbon dioxide gradient in human septic shock. Chest 1992; 101: 509-515
12. Johnson BA, Weil MH. Redefining ischemia due to circulatory failure as dual defects of oxygen deficits and of carbon dioxide excesses. Crit Care Med 1991; 19: 1432-1438
13. Harrison PM, Wendon JA, Gimson AE, Alexander GJ, Williams R. Improvement by acetylcysteine of hemodynamics and oxygen transport in fulminant hepatic failure. N Engl J Med 1991; 324: 1852-1857
14. Wendon JA, Harrison PM, Keays R, Gimson AE, Alexander GJ, Williams R. Effects of vasopressor agents and epoprostenol on systemic hemodynamics and oxygen transport in fulminant hepatic failure. Hepatology 1992; 15: 1067-1071
15. Wendon JA, Harrison PM, Keays R, Gimson AE, Alexander G, Williams R. Arterial-venous pH differences and tissue hypoxia in patients with fulminant hepatic failure. Crit Care Med 1991; 19: 1362-1364
16. Shulman GI, Lacy WW, Liljenquist JE, Keller U, Williams PE, Cherrington AD. Effect of glucose, independent of changes in insulin and glucagon secretion, on alanine metabolism in the conscious dog. J Clin Invest 1980; 65: 496-505
17. Myers SR, Biggers DW, Neal DW, Cherrington AD. Intraportal glucose delivery enhances the effects of hepatic glucose load on net hepatic glucose uptake in vivo. J Clin Invest 1991; 88: 158-167
18. Bratsch-Marrain PR, Waldhäusl WK, Gasić S, Korn A, Nowotny P. Oral glucose tolerance test: effect of different glucose loads on splanchnic carbohydrate and substrate metabolism in healthy man. Metabolism 1980; 29: 289-295
19. Leatherdale BA, Chase RA, Rogers J, Alberti KG, Davies P, Record CO. Forearm glucose uptake in cirrhosis and its relationship to glucose tolerance. Clin Sci (Lond) 1980; 59: 191-198
20. Shangraw RE, Winter R, Hromco J, Robinson ST, Gallaher EJ. Amelioration of lactic acidosis with dichloroacetate during liver transplantation in humans. Anesthesiology 1994; 81: 1127-1138
21. Bernal W, Donaldson N, Wynell D, Wendon J. Blood lactate as an early predictor of outcome in paracetamol-induced acute liver failure: a cohort study. Lancet 2002; 359: 558-563
22. Rolando N, Wade J, Davalos M, Wendon J, Philpott-Howard J, Williams R. The systemic inflammatory response syndrome in acute liver failure. Hepatology 2000; 32: 734-739
23. Salmerón JM, Tito L, Rimola A, Mas A, Navasa MA, Llach J, Ginés A, Ginés P, Arroyo V, Rodés J. Selective intestinal decontamination in the prevention of bacterial infection in patients with acute liver failure. J Hepatol 1992; 14: 280-285
24. Douzinis EE, Tsidemiaidou PD, Pitaridis MT, Andrianakis I, Bobota-Chloraki A, Katsouyanni K, Sfyras D, Malagari K, Roussos C. The regional production of cytokines and lactate in sepsis-related multiple organ failure. Am J Respir Crit Care Med 1997; 155: 53-59
25. Routsi C, Bardouniotou H, Delivoria-Ioannidou V, Kazi D, Roussos C, Zakynthinos S. Pulmonary lactate release in patients with acute lung injury is not attributable to lung tissue hypoxia. Crit Care Med 1999; 27: 2469-2473

S- Editor Guo SY  L- Editor Wang XL  E- Editor Bi L