Temporal Profile of Pneumonia After Stroke

Jeroen C. de Jonge, MD; Diederik van de Beek, MD, PhD; Patrick Lyden, MD; Marian C. Brady, PhD; Philip M. Bath, DSc, FMedSci; H. Bart van der Worp, MD, PhD; on behalf of the VISTA Collaboration*

BACKGROUND AND PURPOSE: The occurrence of pneumonia after stroke is associated with a higher risk of poor outcome or death. We assessed the temporal profile of pneumonia after stroke and its association with poor outcome at several time points to identify the most optimal period for testing pneumonia prevention strategies.

METHODS: We analyzed individual patient data stored in the VISTA (Virtual International Stroke Trials Archive) from randomized acute stroke trials with an inclusion window up to 24 hours after stroke onset and assessed the occurrence of pneumonia in the first 90 days after stroke. Adjusted odds ratios and hazard ratios were calculated for the association between pneumonia and poor outcome and death by means of logistic and Cox proportional hazard regression, respectively, at different times of follow-up.

RESULTS: Of 10,821 patients, 1017 (9.4%) had a total of 1076 pneumonias. Six hundred eighty-nine (64.0%) pneumonias occurred in the first week after stroke. The peak incidence was on the third day and the median time of onset was 4.0 days after stroke (interquartile range, 2–12). The presence of a pneumonia was associated with an increased risk of poor outcome (adjusted odds ratio, 4.8 [95% CI, 3.8–6.1]) or death (adjusted hazard ratio, 4.1 [95% CI, 3.7–4.6]). These associations were present throughout the 90 days of follow-up.

CONCLUSIONS: Two out of 3 pneumonias in the first 3 months after stroke occur in the first week, with a peak incidence on the third day. The most optimal period to assess pneumonia prevention strategies is the first 4 days after stroke. However, pneumonia occurring later was also associated with poor functional outcome or death.

GRAPHIC ABSTRACT: A graphic abstract is available for this article.

Key Words: death ■ pneumonia ■ stroke

About 30% of patients develop an infection in the first days after stroke, of which about one-third is a pneumonia.1 The development of an infection after stroke is associated with a higher risk of poor outcome or death.1–4 In particular, pneumonia contributes to early mortality after stroke, and it has been estimated that 10% of the deaths within 30 days after stroke are attributable to pneumonia.5,6

Limited data from small patients groups suggest that most pneumonias occur in the first 48 to 72 hours after stroke, but these studies only evaluated the development of pneumonia during hospital admission or within the first week or month after stroke.7–10 One study of 369 patients observed 2 peaks in the incidence of pneumonia after stroke: one peak at admission and one peak at day 3 or 4.11 Most of the other previous studies made no further specification on the temporal profile of pneumonia after stroke and this has therefore remained largely unknown.

Greater knowledge about the temporal profile of the relation between pneumonia and functional outcome or death is important to design more effective prevention strategies.12 Knowledge of this temporal profile may identify the most optimal period for (antibiotic) prevention of pneumonia after stroke.
pneumonia after stroke. In 2 large clinical trials on the prevention of infections after stroke, antibiotic treatment was started within 24 or 48 hours of stroke onset, for a period of 4 or 7 days.\textsuperscript{13,14} These trials failed to demonstrate a benefit of preventive antibiotics on the occurrence of pneumonia or on functional outcome. Our objective is to assess the temporal profile of pneumonia in a large number of patients with acute stroke and its relationship with poor outcome or death at different time points.

**METHODS**

We conducted a retrospective analysis of anonymized prospectively collected individual patient data from the acute ischemic stroke or intracerebral hemorrhage databases of the VISTA (Virtual International Stroke Trials Archive). VISTA collects anonymized data from completed randomized stroke trials; its design has been reported previously.\textsuperscript{15} All included studies had individual ethics approval, and all participants gave consent. The data that support the findings of this study are available from the corresponding author upon reasonable request. A completed PRISMA-IPD checklist (Preferred Reporting Items for Systematic Review and Meta-Analyses of Individual Patient Data) is added as Data Supplement.

To assess the occurrence of pneumonia early after stroke onset, we included patient data from trials with an inclusion window up to 24 hours after stroke onset; that reported on the occurrence of pneumonia; and had a follow-up period of 90 days. We expected that all studies would report on the frequency of pneumonia because most pneumonias fulfill the criteria for a serious adverse event (SAE). SAEs that include the term pneumonia on days 0 to 6 and those on days 7 to 90. Other variables of interest included time from stroke onset to start of trial treatment, smoking, and comorbidities (history of diabetes, atrial fibrillation, hypertension, coronary heart disease, heart failure, myocardial infarction, previous stroke or transient ischemic attack, and hypercholesterolemia).

**Statistical Analysis**

The absolute and relative risks of pneumonia or UTI and the median time from stroke onset to onset of pneumonia or UTI were summarized for the first 90 days after stroke. The same numbers were calculated for the first 7 days (day 0–6). The start day of pneumonia was compared in subgroups by means of an independent \( t \) test or Mann-Whitney \( U \) test, where appropriate. Associations between baseline characteristics and the development of pneumonia in the 90 days after stroke were assessed by univariate analysis (\( \chi^2 \) analysis, independent \( t \) test or Mann-Whitney test, where appropriate). Variables with the univariate \( P<0.15 \) were selected for inclusion in a multivariable logistic regression to identify independent predictors of pneumonia. A crude hazard ratio and adjusted hazard ratio with corresponding 95% CI were calculated for the risk of death after pneumonia by means of a Cox proportional hazard model with time-dependent covariates. Crude odds ratios (OR) and adjusted ORs (aOR) with corresponding 95% CIs were calculated for the association between pneumonia and poor outcome by means of logistic regression. In addition, aORs were calculated for (1) the occurrence of pneumonia in specific time periods (days 0–6, days 7–30, days 31–60, and days 61–90) and (2) the date of onset of pneumonia as a continuous variable, in relation to poor outcome and death. The aORs and adjusted hazard ratios were adjusted for factors known to be related to outcome after stroke: age, sex, stroke severity (defined as the score on the NIHSS), stroke type, hypertension, diabetes, myocardial infarction, and treatment with alteplase. Statistical significance was set at 95%. Adjustments for multiplicity of testing were not performed. All data were analyzed using SPSS version 25 (IBM, Chicago, IL).

**RESULTS**

Our study included 10821 patients from both the treatment as placebo group from 9 neutral randomized clinical trials conducted between 1995 and 2013. Table 1 summarizes the baseline characteristics. The mean age was 71 years (interquartile range [IQR], 61–78), 44.3% were female, and the majority of patients had ischemic stroke (89.2%) rather than intracerebral hemorrhage. Since the studies were performed in 1995 and 2013,
none of the patients received mechanical thrombectomy. The median NIHSS score was 12; the study consisted of 234 minor stroke (NIHSS score 0–4), 7017 moderate stroke (NIHSS score 5–15), 2317 moderate to severe stroke (NIHSS score 16–20), and 1251 severe stroke (NIHSS score 21–42).

**Infection Incidence and Temporal Profile**

Thirty-seven pneumonias recorded within 7 days following a previously documented pneumonia were considered a single pneumonia. A total of 1017 patients had a total of 1076 pneumonias in the first 90 days after stroke (cumulative incidence of 9.4%). The median time of pneumonia onset was 4.0 days after stroke (IQR, 2–12). A total of 689 pneumonias (64.0%) occurred in the first 7 days after stroke (Figure 1; cumulative incidence of 6.4%). Of all pneumonias in the first 7 days after stroke, the median time of onset was 3.0 days after stroke (IQR, 2.0–4.0). Most pneumonias were diagnosed on the third day after stroke (Figure 2).

Patients with severe stroke (NIHSS score >12) had a pneumonia earlier (median 4.0 days [IQR, 2–12]) than patients with a less severe stroke (NIHSS score ≤12; median 5.0 days [IQR, 3.0–15.8]; P<0.02). There were no differences in the day of pneumonia onset between subgroups based on age, sex, stroke type, or treatment with alteplase.

In 1033 patients, a total of 1120 UTIs occurred in the first 90 days after stroke (cumulative incidence 9.5%). The median day of UTI onset was 6.0 days (IQR, 3–14 days), which was later than that of pneumonia (P<0.001).

**Predictors of Pneumonia**

Univariate analysis identified higher age, male sex, higher NIHSS, atrial fibrillation, diabetes, coronary heart disease, heart failure, myocardial infarction, and treatment with alteplase as statistically significant predictors of the occurrence of pneumonia in the first 90 days after stroke. In multivariable analysis, higher age, male gender, higher NIHSS, atrial fibrillation, diabetes, and the use of alteplase remained independent predictors (Table 1).

---

**Table 1. Baseline Characteristics**

| Variable                      | All (n=10821) | No pneumonia (n=9804) | Pneumonia (n=1017) | P value |
|-------------------------------|--------------|-----------------------|--------------------|---------|
| Age, y                        | 71 (61–78)   | 71 (61–78)            | 75 (67–81)         | <0.001* |
| Sex (male)                    | 6024/10821 (55.7%) | 5396/9804 (55.0%)  | 628/1017 (61.8%)   | <0.001* |
| NIHSS                         | 12 (9–17)    | 12 (8–17)             | 16 (12–20)         | <0.001* |
| Time from stroke to RCT treat- | 4.2 (3.4–5.4) | 4.2 (3.4–5.5)         | 4.0 (3.3–5.0)      | NA      |
| ment, h                       |              |                       |                    |         |
| Stroke type                   |              |                       | 0.89               |         |
| Ischemic stroke               | 9647/10821 (89.2%) | 8739/9804 (89.1%)   | 908/1017 (89.3%)   |         |
| ICH                           | 1174/10821 (10.8%) | 1065/9804 (10.9%)   | 109/1017 (10.7%)   |         |
| Comorbidity                   |              |                       |                    |         |
| Atrial fibrillation           | 2589/10820 (23.9%) | 2239/9803 (22.8%)   | 350/1017 (34.4%)   | <0.001* |
| Hypertension                  | 7316/10820 (67.8%) | 6600/9803 (67.3%)   | 716/1017 (70.4%)   | 0.046   |
| Diabetes                      | 2269/10821 (21.0%) | 1991/9804 (20.3%)   | 278/1017 (27.3%)   | <0.001* |
| Coronary heart disease        | 2008/7444 (27.0%) | 1737/6596 (26.3%)   | 271/848 (32.0%)    | 0.001†  |
| Heart failure                 | 562/7059 (8.0%) | 481/6262 (7.4%)      | 101/797 (12.7%)    | <0.001† |
| Myocardial infarct            | 1400/10819 (12.9%) | 1244/9802 (12.7%)   | 156/1017 (15.3%)   | 0.017   |
| Previous stroke               | 2879/10809 (26.6%) | 2615/9794 (26.7%)   | 264/1015 (26.0%)   | 0.84    |
| Previous TIA                  | 628/7129 (8.8%) | 552/6320 (8.7%)      | 76/809 (9.4%)      | 0.53    |
| Hypercholesteremia            | 915/4126 (22.2%) | 875/3877 (22.6%)    | 40/249 (16.1%)     | 0.017†  |
| Smoking                       |              |                       | 0.001              |         |
| Never                         | 7813/9978 (78.3%) | 7065/9074 (77.9%)   | 748/904 (82.7%)    | …       |
| Ever                          | 2165/9978 (21.7%) | 2009/9074 (22.1%)   | 156/904 (17.3%)    |         |
| Treatment with alteplase      | 2692/10817 (24.9%) | 2401/9800 (24.5%)   | 291/1017 (28.6%)   | P=0.004* |

Data were presented as median (IQR) or % (n/N). ICH indicates intracerebral hemorrhage; IQR, interquartile range; NIHSS, National Institutes of Health Stroke Scale; RCT, randomized clinical trial; and TIA, transient ischemic attack.

*Remained statistically significant in multivariable analysis.
†Excluded from multivariable analysis because of large number of missing values.
Outcomes
A score on the modified Rankin Scale at 90 days was missing for 241 of the 10,821 patients (2.2%). Of all 1,017 patients with pneumonia after stroke, 485 (47.8%) had died at 90 days, compared with 1,439 (14.7%) of 9,793 patients without pneumonia (P < 0.001; Figure 3). The median time between the diagnosis of pneumonia and death was 6 days [IQR, 2–18]; 8 days [IQR, 3–22] for pneumonia that started between day 0 and 6, and 3 days [IQR, 1–10] for pneumonia that started between day 7 and 90 (P < 0.001). The presence of pneumonia was associated with an increased risk of death (adjusted hazard ratio, 4.1 [95% CI, 3.7–4.6]). The association between pneumonia and a higher risk of death remained present in all analyzed time periods (Table 2). A later start of pneumonia appeared to be associated with a higher risk of death, but this was just not statistically significant (aOR, 1.01 [95% CI, 1.0–1.01]; P = 0.051). Pneumonia in the first 90 days was associated with a poor functional outcome (aOR, 4.8 [95% CI, 3.8–6.1]). The association between pneumonia and poor functional outcome was present in all time periods (Table 2). There was no association between the time of onset of pneumonia and the risk of a poor outcome (aOR, 1.0 [95% CI, 0.99–1.01]; P = 0.74).

DISCUSSION
The current study based on a large international database confirms earlier reports that about one in every 10 patients has a pneumonia in the first 90 days after stroke and shows that almost 2 out of every 3 pneumonias occur in the first week. The peak incidence of pneumonia was on the third day after stroke, accounting for almost 20% of all pneumonias after stroke. Pneumonia occurred earlier and more frequently in patients with a more severe stroke. The occurrence of pneumonia was independently associated with poor functional outcome or death at any time point during the 90-day follow-up period.

Smaller previous studies reported a median or mean period between stroke onset or hospitalization and pneumonia of 1.8, 1.9, 2.0, 3.0, or 4.4 days, observed an peak incidence on the second day, or found that the majority of pneumonias occurred within 48 or 72 hours. However, these studies were limited by a short follow-up duration (eg, only during hospital admission.

Figure 1. Temporal profile of post-pneumonia and urinary tract infection. Absolute number (black bars) and cumulative percentage (gray line) of pneumonia (A) and urinary tract infection (B) in the first 90 d after stroke.
or limited to the first week or month), reported on small patient numbers (with an amount of pneumonias ranging from 22 to 102), combined all infections, or studied specifically patients in the intensive care unit or those with dysphagia or tube feeding.

One study evaluated the course of pneumonia after stroke in more detail. This study evaluated 51 respiratory infections in the first 30 days after stroke onset in 369 tube-fed patients and found 2 peaks of infection (on day 1 and on day 3–4).

In contrast to the previous smaller studies, larger cohort studies on infections or pneumonia after stroke did not provide details on the temporal profile, even emphasizing the limited available data on timing.

In our study, we included 10 821 patients with a follow-up duration of 90 days, in whom we observed 1076 pneumonias. The peak incidence occurred between 48 and 72 hours after stroke onset. The cumulative incidence of 9.4% in our study is comparable to the 10% reported in the literature.

Our finding that pneumonia was associated with poor functional outcome or death at any time point during the 90-day follow-up period is in line with previous findings. In addition, we found that where pneumonia occurred >1 week after stroke onset, it resulted in death in fewer days than when pneumonia occurred during the first week after stroke onset (3 versus 8 days). Our finding may reflect more aggressive pneumonia treatment, dysphagia assessment, or more intensive staff input in stroke units compared with more reserved treatment approaches in a later stage after stroke, for example, because of installed treatment restrictions.

The early peak incidence of pneumonia after stroke in our study is consistent with the 2 most important concepts of the pathophysiology of pneumonia after stroke: stroke-facilitated aspiration and stroke-induced immunosuppression.

In our study, the median time of pneumonia onset was 4.0 days after stroke. However, the moment of diagnosis is preceded by both an incubation period of the pneumonia as well as the time required for diagnostic evaluation to conclude a diagnosis of pneumonia. Therefore, the temporal profile we found suggests that the process towards the development of most pneumonias starts very early after stroke onset. In addition, pneumonias occurred earlier than UTIs, the second most common infection after stroke. Animal studies have suggested that stroke-induced immunosuppression is already present in the first hours after stroke. In addition, about 15% of patients with acute ischemic stroke already have signs of pulmonary infection on chest computed tomography imaging within a few hours after stroke onset, suggestive for aspiration at stroke onset or in the first few hours after stroke. Therefore, the concepts of stroke-facilitated aspiration and stroke-induced immunosuppression are both in accordance with the temporal profile described in the current study.

If preventive antibiotics would be of any benefit, the results of our study suggest that these should be started...
as soon as possible after stroke and continued for at least 4 days. In the large randomized trials, PASS (Preventive Antibiotics in Stroke Study) and STROKE-INF (Prophylactic Antibiotics After Acute Stroke for Reducing Pneumonia in Patients With Dysphagia), prophylactic antibiotics did not improve outcome after 90 days in patients with acute stroke.\textsuperscript{13,14} PASS had an inclusion window of 24 hours and had a treatment duration of 4 days. There was no association between time to start prophylactic treatment (divided in subgroups of 0–3, 3–6, 6–12, and 12–24 hours) and outcome, but the numbers of patients per subgroup are not known. STROKE-INF had an inclusion window of 48 hours after stroke onset and a treatment duration of 7 days. The study did not report on time-to-treatment subgroup analyses. Both studies found that treatment with antibiotics was safe and associated with just a small number of related adverse events, such as allergic reactions or \textit{C difficile} infections. Further insights will be provided by the ongoing PRECIOUS trial (Prevention of Complications to Improve Outcome in Elderly Patients With Acute Stroke), which has a 24-hour inclusion window and includes elderly patients with moderate to severe stroke, who are, therefore, at the highest risk of pneumonia.\textsuperscript{27} Our findings may also inform (randomized trials on) other prevention strategies, such as dysphagia assessment, swallowing interventions, early mobilization, or oral hygiene improvement.

Our study has limitations. First, there is no uniform definition of pneumonia in the VISTA database. Since the date of onset of the pneumonia was the date that was reported on the adverse event logs, the diagnosis was most likely physician-based rather than an made by an expert panel or on the Centers for Disease Control and Prevention criteria.\textsuperscript{28} Previous studies have shown that the incidence of physician-diagnosed pneumonia is higher than that made by an expert panel.\textsuperscript{13,14} In addition, it is not known which sign or symptom was considered by the local investigator to represent the start of pneumonia. Also, in the VISTA database, several terms are used for reporting respiratory tract infections. We

| Time period of pneumonia | Death | aOR (95% CI) | Poor outcome | aOR (95% CI) |
|--------------------------|-------|--------------|--------------|--------------|
| Day 0–6                  | 305/687 (44%) | 3.1 (2.6–3.7) | 618/677 (91%) | 5.0 (3.8–6.7) |
| Day 7–30                 | 122/227 (54%) | 4.6 (3.5–6.2) | 206/225 (92%) | 4.9 (3.0–8.0) |
| Day 31–60                | 57/94 (61%) | 7.5 (4.8–11.8) | 87/94 (93%) | 7.1 (3.1–16.0) |
| Day 61–90                | 29/60 (48%) | 3.8 (2.2–6.6) | 50/58 (86%) | 3.1 (1.4–6.9) |

Frequency of pneumonia in specific time periods and its relation with death and poor outcome, compared with patients without pneumonia. aOR indicates adjusted odds ratio.

Figure 3. Kaplan-Meier curve showing survival of patients with (dark line) and without (gray line) pneumonia after stroke.
have only selected the terms which include the term pneumonia. Terms that could also include other disease entities were not included (eg, lower respiratory tract infection was excluded because acute bronchitis also falls into this category), so the reported pneumonia incidence could be an underestimate. Second, several factors could have influenced the reporting of pneumonia. Since most data on the occurrence of pneumonia are from SAE reporting, there is a risk that some pneumonias could have occurred before inclusion into the trial or that some pneumonias in trials did not meet the criteria for SAE, and therefore, are not incorporated in the current study. We have tried to minimize this risk by selecting a short inclusion window to start of trial treatment of 24 hours. Also, patients who have a pneumonia at baseline, patients who have either the least or most severe stroke, or patients who have severe comorbidities are less likely to be included in a trial, which could have led to selection bias. Third, we do not have information on (antibiotic) treatment of pneumonia. The treatment type, duration of treatment, and the withdrawal of treatment could be important to interpret the relation between pneumonia and death or dependency.

In conclusion, pneumonia is a frequent complication in the first 90 days after stroke, with a peak incidence on the third day. Pneumonia was associated with poor functional outcome or death at any time point during the 90-day follow-up period.

ARTICLE INFORMATION

Received October 5, 2020; final revision received May 30, 2021; accepted June 21, 2021.

The podcast and transcript are available at https://www.wahajournals.org/str/podcast.

Affiliations

Department of Neurology and Neurosurgery, UMC Utrecht Brain Center, University Medical Center Utrecht, Utrecht University, the Netherlands (J.C.d.J., H.B.v.d.W.), Department of Neurology, Amsterdam University Medical Center, Amsterdam Neuroscience, the Netherlands (D.v.d.B.), Departments of Physiology and Neuroscience and Neurology, USC Keck School of Medicine, Los Angeles, CA (P.L.), Nursing, Midwifery and Allied Health Professions Research Unit, Glasgow Caledonian University, United Kingdom (M.C.B.). Stroke Trials Unit, Division of Clinical Neuroscience, School of Medicine, University of Nottingham, United Kingdom (P.M.B.).

Acknowledgments

Dr de Jonge collected data, performed the analyses, and wrote the first draft of the article. All other authors reviewed the article. All authors read and approved the final version of the article.

Sources of Funding

The author(s) disclosed receipt of the following financial support for the research, authorship, and publication of this article: Dr de Jonge is funded by the European Union’s Horizon, 2020 research and innovation programme (grant no. 634809).

Disclosures

Dr van der Worp served as a consultant to Boehringer Ingelheim, Bayer, and LivaNova and received grants from Styrker. Dr van der Worp is Chief Investigator of PRECIOS (Prevention of Complications to Improve Outcome in Elderly Patients With Acute Stroke), a randomized trial assessing the effects of the prevention of complications in patients with acute ischemic stroke. Dr Bath has served on advisory boards with DiaMedica, Moleac, Nestle, Phagogenesis, Platelet Solutions, and Sanofi. Dr Bath is Stroke Association Professor of Stroke Medicine and an Emeritus NIHR Senior Investigator.

SUPPLEMENTAL MATERIALS

Online Appendices I and II

APPENDIX

VISTA Steering Committee: K.R. Lees (Chair), A. Alexandrov, PM. Bath, E. Berge, E. Bhuimki, N. Bornstein, C. Chen, L. Claesson, S.M. Davis, G. Donnan, H.C. Diener, M. Fisher, M. Ginsberg, B. Group, J. Grotta, W. Hacke, M.G. Hennerici, M. Hommel, M. Kaste, P. Lyden, J. Marler, K. Muir, N. Venketasubramanian, R. Sacco, A. Shuaib, P. Teal, N.G. Wahlgren, S. Warach, and C. Weimar.

REFERENCES

1. Westendorp WP, Nederkoorn PJ, Vermeij JD, Dijkgraaf MG, van de Beek D. Post-stroke infection: a systematic review and meta-analysis. BMC Neurol. 2011;11:110. doi: 10.1186/1471-2377-11-110

2. Vermeij FH, Scholle op Reimer WJ, de Man P, van Oostenbrugge RJ, Franke CL, de Jong G, de Kort PL, Dippel DW. Netherlands Stroke Study Inves- tigators. Stroke-associated infection is an independent risk factor for poor outcome after acute ischemic stroke: data from the Netherlands stroke sur- vey. Cerebrovasc Dis. 2009;27:465–471. doi: 10.1159/000210093

3. Finlayson O, Kapral M, Hall R, Asllani E, Selchen D, Saposnik G; Cana- dian Stroke Network; Stroke Outcome Research Canada (SORCan) Working Group. Risk factors, inpatient care, and outcomes of pneumonia after ischemic stroke. Neurology. 2011;77:1338–1345. doi: 10.1212/WNL.0b013e31823152b1

4. Ingeman A, Andersen G, Hundtir HBH, Svendsen ML, Johnsen SP. In-hospital medical complications, length of stay, and mortality among stroke unit patients. Stroke. 2011;42:3214–3218. doi: 10.1161/STROKEAHA.110.601881

5. Katsan IL, Cebul RD, Husak SH, Dawson NV, Baker OW. The effect of pneu- monia on mortality among patients hospitalized for acute stroke. Neurology. 2003;60:620–625. doi: 10.1212/01.wnl.0000046586.38284.60

6. Heuschmann P.U, Kolominsky-Rabas PL, Misselwitz B, Hermanek P, Leffmann C, Janzen RW, Rother J, Buecker-Nott HJ, Berger K; Ger- man Stroke Registers Study Group. Predictors of in-hospital mortality and attributable risks of death after ischemic stroke: the German Stroke Registers Study Group. Arch Intern Med. 2004;164:1761–1768. doi: 10.1001/archinte.164.16.1761

7. Hassan A, Khealani BA, Shafqat S, Aslam M, Salahuddin N, Syed NA, Baig SM, Wasy M. Stroke-associated pneumonia: microbiological data and out- come. Singapore Med J. 2006;47:204–207.

8. Hilker R, Poetter C, Findeisen N, Sobesky J, Jacobs A, Neveling M, Heiss WD. Nosocomial pneumonia after acute stroke: implications for neurological intensive care medicine. Stroke. 2003;34:975–981. doi: 10.1161/01.STR.0000063373.70993.CD

9. Warusavitarane A, Karunatilake D, Sim J, Smith C, Roffe C. Early diag- nosis of pneumonia in severe stroke: clinical features and the diag- nostic role of C-reactive protein. PLoS One. 2016;11:e0150269. doi: 10.1371/journal.pone.0150269

10. Chen LF, Chang CY, Hsu LC, Tsai PH, Chang SJ, Chang SC, Yuan MK, Lai YC, Liu WS. Bacterial pneumonia following acute ischemic stroke. J Chin Med Assoc. 2013;76:78–82. doi: 10.1016/j.jcma.2012.10.005

11. Langdon PC, Lee AH, Bins CW. High incidence of respiratory infections in ‘nil by mouth’ tube-fed acute ischemic stroke patients. Neuroepidemiology. 2009;29:107–113. doi: 10.1159/000177036

12. Learoyd AE, Woodhouse L, Shaw L, Sprigg N, Bereczki D, Berge E, Caso V, Christensen H, Collins R, Czlonkowska A, et al. Infections up to 76 days after stroke increase disability and death. Transl Stroke Res. 2017;8:541– 548. doi: 10.1007/s12975-017-0563-3

13. Westendorp WP, Vermeij JD, Zock E, Hooijenga IJ, Kruyt ND, Bosboom HJ, Kwa VI, Weisfelt M, Remmers MJ, ten Houten M, et al; PASS investigators. The Preventive Antibiotics in Stroke Study (PASS): a pragmatic randomised open-label masked endpoint clinical trial. Lancet. 2015;385:1519–1526. doi: 10.1016/S0140-6736(14)62456-9

14. Kalra L, Irshad S, Hodossi J, Simpson M, Gullford M, Smithard D, Patel A, Rebollo-Mesa I; STROKE-INF Investigators. Prophylactic antibiot- ics after acute stroke for reducing pneumonia in patients with dysphagia (STROKE-INF): a prospective, cluster-randomised, open-label, masked

Reference
endpoint, controlled clinical trial. *Lancet.* 2015;386:1835–1844. doi: 10.1016/S0140-6736(15)00126-9
15. Ali M, Bath P, Brady M, Davis S, Diener HC, Donnan G, Fisher M, Hacke W, Hanley DF, Luby M, et al; VISTA Steering Committees. Development, expansion, and use of a stroke clinical trials resource for novel exploratory analyses. *Int J Stroke.* 2012;7:133–138. doi: 10.1111/j.1747-4949.2011.00735.x
16. de Jonge JC, Tako RAP, Kauw F, de Jong PA, Dankbaar JW, van der Worp HB. Signs of pulmonary infection on admission chest computed tomography are associated with pneumonia or death in patients with acute stroke. *Stroke.* 2020;51:1690–1695. doi: 10.1161/STROKEAHA.120.028972
17. Dziewas R, Ritter M, Schilling M, Konrad C, Delenberg S, Nabavi DG, Stögbauer F, Ringelstein EB, Lüdemann P. Pneumonia in acute stroke patients fed by nasogastric tube. *J Neurol Neurosurg Psychiatry.* 2004;75:852–856. doi: 10.1136/jnnp.2003.019075
18. Brogan E, Langdon C, Brookes K, Blacker D. Can't swallow, can't transfer, can’t toilet: factors predicting infections in the first week post stroke. *J Clin Neurosci.* 2015;22:92–97. doi: 10.1016/j.jocn.2014.05.035
19. Sellars C, Bowie L, Bagd J, Sweeney MP, Miller H, Tilton J, Langhorne P, Stott DJ. Risk factors for chest infection in acute stroke: a prospective cohort study. *Stroke.* 2007;38:2284–2291. doi: 10.1161/STROKEAHA.106.478156
20. Matz K, Seyfang L, Dachenhansen A, Teuschl Y, Brainin M; MD for the Austrian Stroke Unit Registry Collaborators. Post-stroke pneumonia at the stroke unit - a registry based analysis of contributing and protective factors. *BMC Neurosci.* 2016;17:107. doi: 10.1186/s12888-016-0627-y
21. Kishore AK, Vai L, Chamorro A, Garau J, Hopkins SJ, Di Napoli M, Kalaria L, Langhorne P, Montaner J, Roffe C, et al. How is pneumonia diagnosed in clinical stroke research? A systematic review and meta-analysis. *Stroke.* 2015;46:1202–1209. doi: 10.1161/STROKEAHA.114.007843
22. Ullm L, Harms H, Ohiraun S, Reimnitz P, Meisel A. Impact of infections on long-term outcome after severe middle cerebral artery infarction. *J Neurol.* 2012;319:15–17. doi: 10.1016/j.jnr.2012.05.042
23. Bray BD, Smith CJ, Cloud GC, Enderby P, James M, Paley L, Tyrrell R, Wolfe CD, Rudd AG; SSNAP Collaboration. The association between delays in screening for and assessing dysphagia after acute stroke, and the risk of stroke-associated pneumonia. *J Neurol Neurosurg Psychiatry.* 2017;88:25–30. doi: 10.1136/jnnp-2016-313356
24. Kumar S, Selim MH, Caplan LR. Medical complications after stroke. *Lancet Neurol.* 2010;9:105–118. doi: 10.1016/S1474-4422(09)70266-2
25. Chamorro A, Meisel A, Planas AM, Urra X, van de Beek D, Veltkamp R. The immunology of acute stroke. *Nat Rev Neurol.* 2012;8:401–410. doi: 10.1038/nrneurol.2012.98
26. Prass K, Meisel C, Höflisch C, Braun J, Haller E, Wolf T, Ruscher K, Victorov IV, Plitter J, Dimagli U, et al. Stroke-induced immunodeficiency promotes spontaneous bacterial infections and is mediated by sympathetic activation reversal by poststroke TH1 helper cell type 1-like immunostimulation. *J Exp Med.* 2003;198:725–736. doi: 10.1084/jem.20021098
27. Reinink H, de Jonge JC, Bath PM, van de Beek D, Berge E, Borregaard S, Ciccone A, Csiba L, Demotes J, Dippel DW, et al. PRECIOUS: prevention of complications to improve outcome in elderly patients with acute stroke. Rationale and design of a randomised, open, phase III, clinical trial with blinded outcome assessment. *Eur Stroke J.* 2018;3:291–298. doi: 10.1177/239698731772687
28. Smith CJ, Kishore AK, Vai L, Chamorro A, Garau J, Hopkins SJ, Di Napoli M, Kalaria L, Langhorne P, Montaner J, et al. Diagnosis of stroke-associated pneumonia: recommendations from the pneumonia in stroke consensus group. *Stroke.* 2015;46:2335–2340. doi: 10.1161/STROKEAHA.115.009617