NT-proBNP (N-Terminal Pro-B-Type Natriuretic Peptide) and the Risk of Stroke

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NT-proBNP (N-Terminal Pro-B-Type Natriuretic Peptide) and the Risk of Stroke
Results From the BiomarCaRE Consortium

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Background and Purpose—NT-proBNP (N-terminal pro-B-type natriuretic peptide) is a risk factor for atrial fibrillation and a marker of cardiac function used in the detection of heart failure. Given the link between cardiac dysfunction and stroke, NT-proBNP is a candidate marker of stroke risk. Our aim was to evaluate the association of NT-proBNP with stroke and to determine the predictive value beyond a panel of established risk factors.

Methods—Based on the Biomarkers for Cardiovascular Risk Assessment in Europe-Consortium, we analyzed data of 58,173 participants (50% men; mean age 52 y) free of stroke from 6 community-based cohorts. NT-proBNP measurements were performed in the central Biomarkers for Cardiovascular Risk Assessment in Europe laboratory. The outcomes considered were total stroke and subtypes of stroke (ischemic/hemorrhagic).

Results—During a median follow-up time of 7.9 years, we observed 1,550 stroke events (1,176 ischemic). Increasing quartiles of the NT-proBNP distribution were associated with increasing risk of stroke ($P$ for trend <0.0001; multivariable Cox regression analysis adjusted for risk factors and cardiac diseases). Individuals in the highest NT-proBNP quarter (NT-proBNP $>82.2$ pg/mL) had 2-fold (95% CI, 75%–151%) greater risk of stroke than individuals in the lowest quarter (NT-proBNP $<20.4$ pg/mL). The association remained unchanged when adjusted for interim coronary events during follow-up, and though it was somewhat heterogeneous across cohorts, it was highly homogenous according to cardiovascular risk profile or subtypes of stroke. The addition of NT-proBNP to a reference model increased the C-index discrimination measure by 0.006 ($P=0.0005$), yielded a categorical net reclassification improvement of 2.0% in events and 1.4% in nonevents and an integrated discrimination improvement of 0.007.

Conclusions—In European individuals free of stroke, levels of NT-proBNP are positively associated with risk of ischemic and hemorrhagic stroke, independently from several other risk factors and conditions. The addition of NT-proBNP to variables of established risk scores improves prediction of stroke, with a medium effect size. (Stroke. 2019;50:00-00. DOI: 10.1161/STROKEAHA.118.023218.)

Key Words: atrial fibrillation ■ biomarkers ■ brain ■ epidemiology ■ stroke

NT-proBNP (N-terminal pro-B-type natriuretic peptide) is the N-terminal fragment of the B-type natriuretic peptide, secreted by myocytes as a reaction to several stimuli including wall stretch. NT-proBNP has a central role in the regulation of blood pressure, blood volume, and sodium balance. Its levels increase with age, ventricular hypertrophy and in acute coronary syndromes, heart failure, and atrial fibrillation. NT-proBNP is considered a valuable predictor in diagnosis and prognosis of patients with symptoms of heart failure, left ventricular dysfunction, and acute coronary
syndromes. Several studies have investigated the association of NT-proBNP with an occurrence of cardiovascular or stroke events in general populations. A meta-analysis of 40 prospective studies has demonstrated a clear association between high levels of NT-proBNP and increased cardiovascular risk under a range of different conditions but had insufficient power to assess whether NT-proBNP was associated differently with ischemic or hemorrhagic stroke or with fatal or nonfatal stroke. In the ARIC study (The Atherosclerosis Risk in Communities Study), NT-proBNP was found to be associated with total stroke, nonlacunar ischemic, and especially cardioembolic stroke but not with lacunar or hemorrhagic stroke. In a case-cohort study derived from the REGARDS cohort (Reasons for Geographic and Racial Differences in Stroke), the authors confirmed that the association of NT-proBNP with stroke was largest for cardioembolic stroke.

Using the harmonized database and biobank of the Biomarkers for Cardiovascular Risk Assessment in Europe (BiomarCaRE) project (FP7/2007–2013), we centrally analyzed individual NT-proBNP concentrations in 40,336 individuals of 6 population cohorts with the aims: to achieve a precise characterization of the association of NT-proBNP with stroke in Europe; to assess possible difference in the association of NT-proBNP with ischemic or hemorrhagic stroke or with fatal stroke; and to determine the predictive value of NT-proBNP beyond classical risk factors for stroke.

Methods

Study Overview

Data, analytic methods, and study materials are not available to other researchers.

The present analysis is based on data from the BiomarCaRE consortium (http://www.biomarcare.eu), details of which have been described previously. BiomarCaRE is based on the MONICA Risk Genetics Archiving and Monograph (MORGAM) Project. The MORGAM/BiomarCaRE Data Center in Helsinki harmonized individual data from 15 population-based cohort studies with central storage of selected biological samples of almost 187,736 participants in the BiomarCaRE laboratory at the University Heart Center, Hamburg. Current analyses include cohorts with available information on stroke status at baseline, with adjudication for stroke at follow-up and available data on NT-proBNP (n=6 cohort studies). All NT-proBNP levels included in the present study were measured centrally using the same assay. Statistical analyses were planned and conducted at the NEUROMED BiomarCaRE Center in Pozzilli, Italy. Our study complies with the Declaration of Helsinki, all participating studies were approved by local ethics review boards and written informed consent was obtained from individuals.

Study Cohorts

Overall, the cohort consisted of 6 population-based studies involving 58,173 individuals free of stroke at baseline (individuals with a positive history of stroke based on self-report or prior physician’s diagnosis for stroke were excluded from analyses). The individual cohorts were FINRISK 1997, MONICA Northern Sweden, Prospective Epidemiological Study of Myocardial Infarction from Belfast (PRIME), MATISS Rome, MONICA Brienza, and Moli-sani. Each cohort is based on representative population samples. Full details of baseline data have been provided elsewhere. Cohort descriptions are provided in Table I in the online-only Data Supplement. For each cohort the following harmonized variables were available at baseline: age, sex, body mass index, systolic and diastolic blood pressure, antihypertensive medication, smoking status, total and HDL (high-density lipoprotein) cholesterol and estimated glomerular filtration rate calculated by using of Chronic Kidney Disease Epidemiology Collaboration formula, history of diabetes mellitus, myocardial infarction, atrial fibrillation, and heart failure. Data on atrial fibrillation or heart failure were not available for MATISS and MONICA Brienza. History of diabetes mellitus was defined as self-reported or documented diabetes mellitus. Data collection on risk factors followed a standardized protocol described in the MORGAM Manual.

Laboratory Procedures

NT-proBNP levels were measured in the BiomarCaRE core laboratory using an electrochemiluminescence sandwich immunosay (ECLIA, Roche Diagnostics) on either the ELECSYS 2010 or the Cobas e411 system. The analytical range was 5 to 35,000 pg/mL. The study-specific intra-assay and interassay coefficients of variation are described in Table II in the online-only Data Supplement.

Study Outcome

Participants in each cohort were followed-up for first stroke (fatal or nonfatal) and death from other causes. Deaths were identified through record linkage with national or regional health information systems. Nonfatal strokes refer to survival at 28th day after onset and were identified by linkage to population registers, hospital discharge data, or direct contact with the participant. Most centers adjudicated the events using MONICA diagnostic criteria. The MORGAM Manual gives further information about the event classifications. The procedure used for identification of stroke subtypes (ischemic or hemorrhagic) is described in details in Methods in the online-only Data Supplement. Briefly, a stroke is classified as a cerebral infarction (ischemic stroke) if at least one of the following is present: (1) validation of recent brain infarction in necropsy; (2) circumscribed hypodensity changes of recent origin in the brain parenchyma on computed tomography (CT); and (3) typical signs of infarct in the brain parenchyma on magnetic resonance imaging. The event was considered as cerebral infarction also if there was no validation as described above but the routine clinical or causes of death diagnoses indicated cerebral infarction (International Classification of Diseases, Eighth Revision [ICD-8] value of 432, 433, or 434; an International Classification of Diseases, Ninth Revision [ICD-9] value of 434; or an International Classification of Diseases, Tenth Revision [ICD-10] value of I63). To be accepted as a case of subarachnoid hemorrhage, at least one of the following must be present: (1) necropsy—recent subarachnoid hemorrhage; (2) CT-signs of blood in the subarachnoid cisterns or in cerebral ventricles; (3) magnetic resonance imaging—signs of blood in the subarachnoid cisterns or in cerebral ventricles; and (4) cerebrospinal fluid (liquor) bloody and xanthochromic and the possibility of intracerebral hemorrhage excluded by necropsy or CT examination.

To be accepted as a case of intracerebral hemorrhage in MORGAM, at least one of the following must be present: (1) necropsy—recent intracerebral hemorrhage; (2) CT-hyperdensity changes in the brain parenchyma; (3) magnetic resonance imaging—typical signs of bleeding in the brain parenchyma; and (4) cerebrospinal fluid (liquor) bloody in the presence of focal neurological signs at onset. The event was considered as hemorrhagic stroke also if there was no validation for subarachnoid or intracerebral hemorrhage as described above but the routine clinically recorded or officially registered causes of death diagnoses indicated hemorrhagic stroke (ICD-8 value of 430 or 431, an ICD-9 value of 430 or 431, or an ICD-10 value of I60 or I61).

Statistical Analysis

N=4054 (7.0%) individuals had NT-proBNP values below the limit of detection (5 pg/mL); for these individuals NT-proBNP values have been imputed to NT-proBNP=5 pg/mL. For 9.0% of the available population, one or more cardiovascular disease (CVD) risk factors or NT-proBNP levels were missing; in these cases, we used multiple imputation techniques (SAS PROC MI, n=10 imputed datasets; and PROC MIANALYZE) to maximize data availability.

The NT-proBNP distribution in the overall cohort was right-skewed (mean 82 pg/mL, SD 232 pg/mL; median 43 pg/mL.
coefficient of skewness 35.8). After a natural log transformation, the NT-proBNP distribution showed a Gaussian distribution (mean 3.7 log [pg/mL], SD 1.1 log [pg/mL]; median 3.7 log [pg/mL], coefficient of skewness 0.12). Hereafter, the natural log of NT-proBNP levels has been used. The correlation between log(NT-proBNP) and sex, examination age, total and HDL cholesterol, smoker status, hypertension, systolic and diastolic blood pressure, body mass index, diabetes mellitus, and estimated glomerular filtration rate was assessed using Pearson coefficient. To estimate the association between NT-proBNP and stroke outcome, we first derived sample quartiles for the marker in the pooled sample. Actually, because the log transformation is a monotonic transformation, quartiles constructed by using log(NT-proBNP) values or by NT-proBNP values are identical. More importantly, the creation of quarters and the values of the quartiles are independent from the method of imputation for values under the limit of detection. Subsequently, we estimated the hazard ratios (HRs; with 95% CI) for stroke across increasing NT-proBNP quartiles from Cox proportional hazards models with age as the time scale, and adjusting for sex, study center (Model 1), smoking, body mass index, diabetes mellitus, myocardial infarction at baseline, hypertension medication, total and HDL cholesterol (Model 2). We selected possible confounding variables for regression models based on previous analyses from the same populations. Additional adjustment was also made for baseline estimated glomerular filtration rate or for coronary heart disease, atrial fibrillation or heart failure as time-dependent variables, as these events occurred during follow-up (Model 3). From this latter analysis, the MATISS Study and MONICA Brianza Study were excluded because data on atrial fibrillation or heart failure were not available. Associations of NT-proBNP with each stroke subtype were estimated after censoring participants when they developed stroke of another subtype. We reported the Cochran Q test and the I² statistic estimated after censoring participants when they developed stroke of another subtype. We tested the Cochran Q test and the P statistic to quantify heterogeneity among cohorts. The C-index, the categorical net reclassification improvement and the absolute and relative information to determine the type of stroke (Table 1).

Results

Baseline Characteristics

Baseline characteristics for the overall study population are shown in Table 1. The sex ratio of the overall cohort was balanced with 50% males. The median age was 52 years (interquartile range 43–61 years). Study participants were slightly overweight (median body mass index 27.3 kg/m²). At baseline 24.7% of the individuals were daily smokers, 19.2% were prescribed antihypertensive medication, 5% had diabetes mellitus, and <2.7% had a personal history of myocardial infarction, heart failure, or atrial fibrillation (Table 1). Characteristics of the different cohorts are illustrated in Table III in the online-only Data Supplement.

NT-proBNP levels were lower in men (r=−0.24) and correlated positively with age (r=0.45). The correlation of log(NT-proBNP) levels with other cardiovascular risk factors and phenotypes was generally modest (Table 2).

We found slightly higher (+0.069, SD=0.008) log(NT-proBNP) age and sex-adjusted levels in the northern Europe cohorts (Finland, Sweden, and UK-Belfast) compared with the southern Europe cohorts (Italian cohorts; P<0.0001).

Table 1. General Characteristics of the Studied Population

<table>
<thead>
<tr>
<th>Baseline Characteristics</th>
<th>No. of populations, N</th>
<th>No. of individuals, N</th>
<th>Years of baseline examinations, range in years</th>
<th>Age at baseline examination, y</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>6</td>
<td>58 173</td>
<td>1986–2008</td>
<td>52 (13)</td>
</tr>
<tr>
<td>Men, N (%)</td>
<td>29 275 (50.32)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Myocardial infarction, N (%)</td>
<td>1571 (2.70)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Heart failure, N (%)</td>
<td>505/49 009 (1.03)*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Atrial fibrillation, N (%)</td>
<td>523/42 532 (1.23)*</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

End points during follow-up

| Stroke (any type), N (%) | 1550 (2.66) |
| Ischemic stroke, N (%)† | 1176 (2.02) |
| Hemorrhagic stroke, N (%)† | 330 (0.57) |
| Fatal stroke (any type), N (%) | 249 (0.43) |

Other events during follow-up

| Myocardial infarction, N (%) | 776 (1.33) |
| Heart failure, N (%)‡ | 1431/46 012 (3.11)‡ |
| Atrial fibrillation, N (%)‡ | 1363/45 775 (2.98)‡ |

Characteristics are presented as absolute and relative frequencies for categorical variables, and mean value and SD for continuous variables as well as ranges in years for years of baseline examinations. HDL indicates high-density lipoprotein; and NT-proBNP, N-Terminal Pro-B-type natriuretic peptide.

†Incidence of heart failure at baseline was available for 49 009 individuals. History of atrial fibrillation at baseline was available for 42 532 individuals.

‡Incidence of heart failure (atrial fibrillation) was not evaluated for 12 161 (21.3%) individuals.

NT-proBNP Concentrations and Association With Stroke Outcomes

During a median follow-up time of 7.9 years (interquartile range 4.2–13.8) n=1550 incident stroke events (n=249 fatal events) occurred. Of these, n=1176 were ischemic, n=330 were hemorrhagic (35% of them were subarachnoid and 65% intracerebral hemorrhages), and for n=44 there was insufficient information to determine the type of stroke (Table 1).

Table 3 displays fully adjusted HRs across quarters of the NT-proBNP distribution indicating the associations with
stroke. The risk of stroke in the top quarter was double that in the bottom quarter. The association was virtually unchanged when adjusted for estimated glomerular filtration rate and it was slightly attenuated when further adjusted for occurrence of coronary events or atrial fibrillation or heart failure during follow-up (before the time of stroke event, for an individual who had a stroke; Table 3, Model 3). These associations were of similar magnitude for both ischemic and hemorrhagic and for fatal and nonfatal strokes (Table 3).

The adjusted HR (Model 2) for total, ischemic, or hemorrhagic stroke associated with one SD increase of log(NT-proBNP) concentration was 1.48 (95% CI, 1.40–1.57), 1.51 (95% CI, 1.41–1.61), and 1.45 (95% CI, 1.27–1.66), respectively.

The adjusted HR (Model 2) for case fatality (stroke fatal events among stroke events) for 4° versus 1° quarter of NT-proBNP was 1.16 (95% CI, 0.71–1.92).

Subgroup Analysis of the NT-proBNP-Associated Risk
The distribution of stroke events in quarters of NT-proBNP across cohorts is reported in Table III in the online-only Data Supplement. Figures 1 and 2 display the adjusted HR (Model 2 as defined in Table 3) for total stroke for 4° versus 1° quarter of NT-proBNP across cohorts (Figure 1) and in subgroups of individuals with different cardiovascular risk profiles (Figure 2). The association of NT-proBNP with increased risk of total stroke was observed in all cohorts, with a negligible level of heterogeneity (Figure 1; Cochran Q=3.07, P=0.69; I²=0%).

The effect of NT-proBNP was highly homogeneous across CVD risk categories (Figure 2), with the exception of HDL (the association of NT-proBNP with stroke was greater in individuals with HDL >53 mg/dL). The relative risk of stroke for 4° versus 1° quarter of NT-proBNP was also similar in individuals free from CVD at baseline (myocardial infarction, atrial fibrillation, or heart failure, n=40,507, n=1100 stroke events during follow-up), HR=2.21 (95% CI, 1.79–2.72) when compared with individuals who did report a history of myocardial infarction or atrial fibrillation or heart failure, n=2313, n=212 stroke events at baseline, HR=2.15 (95% CI, 0.93–4.99; P for difference =0.73).

NT-proBNP and Prediction of Stroke
The addition of NT-proBNP to the base model increased the C-index discrimination measure by 0.006 (from 0.842 to 0.848; P value for testing increment equal to zero: 0.0005), yielded a net reclassification improvement of 2.0% in events and 1.4% in nonevents, and an absolute and relative integrated discrimination improvement of 0.007 and 0.11, respectively.

Discussion
Based on harmonized individual-level data and a centrally standardized NT-proBNP evaluation in >58,000 individuals from 6 population-based European studies, our analyses indicate that high levels of NT-proBNP (in particular in the upper quarter of the distribution, >82.2 pg/mL) are a risk factor for stroke, independent of conventional risk factors. This supports growing evidence associating high levels of natriuretic peptides with increased risk of stroke.

Comparison to Previous Studies
Our findings are in agreement with a meta-analysis of 13 studies including 2063 stroke events in 56,764 individuals which found a relative risk of 1.93 (95% CI, 1.58–2.37) among individuals in the top third in comparison to the bottom third of the NT-proBNP distribution. We also confirmed the association of NT-proBNP with ischemic stroke, as observed in the REGARDS study. Unfortunately, we were unable to distinguish cardioembolic strokes and consequently cannot confirm the interesting findings of both the REGARDS study and the ARIC study which found that the associations of NT-proBNP with stroke events were strongest for cardioembolic strokes.

NT-proBNP as a Stroke Risk Factor
We found slightly higher NT-proBNP levels in populations from northern Europe compared with those in southern Europe. However, the difference represents only 6.3% of the SD of the peptide distribution.

The association of NT-proBNP levels with stroke was absent when the second quarter was compared with the first, modest when the third quarter was compared with the first and evident when the top quarter is compared with the bottom. Our findings indicate that the critical value above which the risk of stroke becomes important is around NT-proBNP=80 pg/mL. This value accords well with the corresponding threshold observed in the ARIC study (80 pg/mL) though it is lower than that observed in the REGARDS study (137 pg/mL).

Interestingly, we observed an association of NT-proBNP with incidence of both ischemic and hemorrhagic stroke and with both fatal and nonfatal stroke. A link of NT-proBNP with risk of ischemic stroke is not unexpected given the correlation of NT-proBNP with cardiac function. To our knowledge, this is the first observation of a statistically significant association of high levels of NT-proBNP with risk of hemorrhagic stroke. In the ARIC study, higher levels of NT-proBNP were
associated with an almost double risk of hemorrhagic stroke, but the small number of events (n=63) made the observation statistically imprecise. The association of NT-proBNP with different types of stroke may be because of shared risk factors, unidentified effects of NT-proBNP or unknown mechanisms for these strokes. Plasma brain natriuretic peptide levels have been shown to be elevated not only in acute ischemic stroke patients but also in the acute phase of subarachnoid and intracerebral hemorrhage. Interestingly, growing evidence suggests causal relationships of natriuretic peptides to endothelial permeability, which might predispose not only to atherosclerosis, but to hemorrhages too. In fact, Lee et al demonstrated that salt-loaded stroke-prone spontaneous hypertensive rats have increased vascular permeability at the site of subsequent intracerebral hemorrhage, and Lin et al demonstrated that elevated permeability predicted subsequent hemorrhagic transformation following ischemic stroke.

The stroke risk associated with elevated NT-proBNP levels is highly homogenous according to the presence or absence of other cardiovascular risk factors, suggesting that raised NT-proBNP levels affect risk for stroke over a broad spectrum of circumstances, and in particular, independent of the presence of hypertension.

The role of NT-proBNP as risk factor for stroke was comparable in individuals with or without cardiac diseases at baseline. In addition, adjustment for the presence of CVD at baseline and during follow-up before the stroke event only slightly modified the association between NT-proBNP and total or ischemic stroke. These findings suggest that the association between NT-proBNP and stroke is not secondary to the occurrence of other CVD that could associate with both NT-proBNP levels and stroke.

NT-proBNP is mainly released by cardiac myocytes and is only weakly associated with other CVD risk factors, except age and sex. In accord with the previous finding of a specific association with cardioembolic stroke, elevated NT-proBNP at baseline is the most probably because of subclinical cardiac pathology which increases the risk of stroke events years later. In this case, an elevated NT-proBNP should

| Table 3. Hazard Ratios for Different Stroke Outcomes, According to Quarters of NT-ProBNP |
|---------------------------------|---------------------------------|---------------------------------|---------------------------------|---------------------------------|
|                                | NT-ProBNP Quarters (pg/mL)      |                                |                                |                                |
|                                | <20.4                          | 20.4–42.5                      | 42.5–82.2                      | >82.2                          |
|                                | N                              | 14,543                         | 14,545                         | 14,541                         | 14,544                         |
| All strokes (N=1550)           | No. of events                  | 207                            | 248                            | 321                            | 774                            |
| Model 1* (95% CI)              | -1- reference                  | 1.09 (0.89–1.32)               | 1.27 (1.05–1.53)               | 2.25 (1.88–2.68)               |
| Model 2† (95% CI)              | -1- reference                  | 1.09 (0.90–1.32)               | 1.28 (1.06–1.54)               | 2.14 (1.79–2.57)               |
| Model 3‡ (95% CI)              | -1- reference                  | 1.09 (0.90–1.32)               | 1.27 (1.05–1.53)               | 2.06 (1.72–2.47)               |
| Ischemic strokes (N=1176)      | No. of events                  | 152                            | 189                            | 235                            | 600                            |
| Model 2† (95% CI)              | -1- reference                  | 1.13 (0.91–1.42)               | 1.28 (1.02–1.60)               | 2.24 (1.82–2.76)               |
| Model 3‡ (95% CI)              | -1- reference                  | 1.12 (0.90–1.40)               | 1.25 (1.00–1.57)               | 2.12 (1.72–2.62)               |
| Hemorrhagic strokes (N=330)    | No. of events                  | 50                             | 53                             | 76                             | 151                            |
| Model 2† (95% CI)              | -1- reference                  | 1.01 (0.67–1.51)               | 1.41 (0.96–2.06)               | 2.09 (1.42–3.09)               |
| Model 3‡ (95% CI)              | -1- reference                  | 1.02 (0.68–1.53)               | 1.42 (0.97–2.08)               | 2.09 (1.42–3.09)               |
| Incident strokes that were fatal (N=249) | No. of events                  | 25                             | 34                             | 48                             | 142                            |
| Model 2† (95% CI)              | -1- reference                  | 1.08 (0.63–1.84)               | 1.09 (0.64–1.86)               | 2.15 (1.33–3.47)               |
| Model 3‡ (95% CI)              | -1- reference                  | 1.11 (0.65–1.88)               | 1.12 (0.66–1.89)               | 2.08 (1.29–3.37)               |
| Incident strokes that were nonfatal (N=1301) | No. of events                  | 182                            | 214                            | 273                            | 632                            |
| Model 2† (95% CI)              | -1- reference                  | 1.10 (0.89–1.36)               | 1.32 (1.08–1.62)               | 2.15 (1.77–2.62)               |
| Model 3‡ (95% CI)              | -1- reference                  | 1.10 (0.89–1.35)               | 1.31 (1.07–1.61)               | 2.07 (1.70–2.52)               |

BMI indicates body mass index; HDL, high-density lipoprotein; and NT-proBNP, N-Terminal Pro-B-type natriuretic peptide.

*Model 1: adjusted for age, sex, and center.
†Model 2: model 1 + smoking, BMI, diabetes mellitus, hypertension medication, systolic and diastolic blood pressure, total and HDL cholesterol, myocardial infarction at baseline.
‡Model 3: model 2 + coronary heart disease, atrial fibrillation or heart failure as time-dependent variables as these events occurred during follow-up.
prompt careful considerations and diagnosis of potential underlying cardiac problems, which if treated appropriately, may prevent future adverse events.

The addition of NT-proBNP on top of several stroke risk factors improved both discrimination and reclassification. The magnitude of improvement was comparable to that of troponin for coronary heart disease, as demonstrated in similar large collaborative studies of population-based cohorts. In addition, the relative integrated discrimination improvement indicates that the strength of NT-proBNP is larger than the average strength of risk factors in the reference model, according to the criterion suggested by Pencina et al. Therefore, all considered, the effect size of NT-proBNP for stroke prediction in the general population can be considered to range from moderate to medium.

Limitations
Some strengths and limitations of the present study should be considered. Although our validation of stroke events was systematic and detailed, it was based, as is usually the case in most epidemiological studies, on medical reviews and not on standardized neurological examinations or data from CT or magnetic resonance imaging, especially for those cohorts with baseline enrollment in 1980s and 1990s. Moreover, information on the cause of ischemic strokes, such as the presence of a cardiac source of embolism, would have been valuable for the analysis but unfortunately was not generally available. Because of the low number of hemorrhagic strokes we decided to not conduct separate analyses for subarachnoid or intracerebral hemorrhages. Despite longstanding expertise in data harmonization in the MORGAM Data Centre in Helsinki since 1998, resulting in the best possible end point and covariate validation, measurement error and lack of information on some other known cardio-metabolic risk factors (such as physical activity and diet), offers some room for residual confounding to affect the observed associations among the more than 58,000 individuals investigated in these 6 European population-based cohort studies. On one hand, we present a large dataset of NT-proBNP values measured centrally with
the same assay, but on the other, the differences in storage duration among the included cohorts may have led to differences in variability in NT-proBNP levels across cohorts, but given the broad homogeneity of the relationship with stroke risk, we see no reason why this would bias the observed associations. Further, since we had only single measurements of NT-proBNP, we cannot correct for regression dilution bias. This could have led to an underestimation of our risk estimates. We also cannot examine, with a single measure, how risk of stroke might vary when biomarker levels change over time.

Conclusions
In this the largest transnational dataset with centrally measured NT-proBNP, we confirm NT-proBNP as a risk factor for ischemic stroke and also demonstrate an association with hemorrhagic stroke. NT-proBNP measurement might support the identification of those individuals at high risk for stroke, who would benefit most from preventive interventions.

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Disclosures
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