Stroke mimics in patients clinically diagnosed with stroke at a tertiary teaching hospital in Tanzania: a prospective cohort study

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Abstract

Background: Stroke mimics account for up to one-third of acute stroke admissions and are a heterogeneous entity which pose diagnostic challenges. Diagnosing such patients is however crucial to avoid delays in treatment and potentially harmful medication prescription. We aimed at describing the magnitude, clinical characteristics and short-term outcomes of stroke mimics in patients clinically diagnosed with a stroke.

Methods: This prospective study enrolled patients admitted with a World Health Organization clinical criteria for stroke at a tertiary hospital in Tanzania. Baseline data was collected and the simplified version of the FABS scale was used to determine its usefulness in predicting stroke mimics. The National Institute of Health Stroke Scale and Modified Rankin Scale were used to assess for admission stroke severity and outcomes respectively.

Results: Among 363 patients with suspected stroke on admission, the final diagnosis was stroke mimics in 24 (6.6%) who had a mean age of 65.8 ± 15 years. Patients with stroke mimics were less likely to have cardiovascular risk factors for stroke including premorbid hypertension (7 (29.2%) vs 263 (77.6%), p < 0.001) and increased waist-hip ratio (9 (37.5%) vs 270 (79.6%); p < 0.001) for mimics and true strokes respectively. Clinical findings such as hypertension and the presence of cortical features in neurological examination occurred less in patients with stroke mimics. The simplified FABS score of ≥ 3 could identify patients with stroke mimics with a sensitivity and specificity of 38 and 80% respectively. The most common causes of mimics were brain tumors 6 (25%), meningoencephalitis 4 (16.7%) and epileptic seizures 3 (12.5%). The majority of patients with stroke mimics had severe disease on admission and the 30-day mortality in these patients was 54.5%.

Conclusions: In the present study, the proportion of stroke mimics among patients clinically diagnosed with stroke was 6.6% and brain tumors was a common etiology. Stroke mimics were less likely to have cardiovascular risk factors and cortical signs during evaluation. We recommend further studies that can help develop clinical scales used for predicting stroke mimics in an African population.

Keywords: Stroke, Mimics, Magnitude, Mortality, Outcomes

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Background

Stroke is a clinical diagnosis as endorsed by World Health Organization (WHO), defined as rapidly developing clinical signs of focal or global disturbance in cerebral function lasting more than 24 h or leading to death with no apparent cause other than that of vascular origin [1]. It is a medical emergency and an early diagnosis is of paramount importance. This entails judicious clinical assessment supported with neuroimaging studies to confirm the diagnosis. Certain conditions however can imitate true strokes, leading to a misdiagnosis, delays and/or inappropriate treatment. Such disorders are termed stroke mimics (SM); pathological conditions that resemble stroke like clinical presentations, however are caused by diseases other than cerebrovascular disorders [2]. In modern era where thrombolytics are ever are caused by diseases other than cerebrovascular conditions however can imitate true strokes, leading to a misdiagnosis, delays and/or inappropriate treatment. Such disorders are termed stroke mimics (SM); pathological conditions that resemble stroke like clinical presentations, however are caused by diseases other than cerebrovascular disorders [2]. In modern era where thrombolytics are widely used in treating acute stroke, evidence indicates that as many as a 15% of patients treated with tissue plasminogen activator (t-PA) are SM for which the therapy is not indicated [3]. This results in unnecessary diagnostic tests, invasive procedures and longer hospital stay leading to an increased cost to the patient.

Previous data from high income states describes the proportion of SM as high as 30% of all strokes [4]. Similarly, in sub Saharan Africa (SSA), a hospital based retrospective study conducted in Nigeria, found that among 142 patients with a WHO clinical diagnosis of stroke, 23.2% had SM which included brain tumors, brain abscess, subdural hematomas, hydrocephalus and intracranial cysts [5]. Other described SM include seizures, syncope, metabolic encephalopathies (such as hypoglycemia, hyponatremia, uremia and hepatic encephalopathy), infectious disorders including meningoencephalitis and degenerative diseases amongst others [6, 7]. Notable is that, in SSA, stroke is readily diagnosed purely on clinical grounds with no confirmation due to limited access to brain imaging [8]. Therefore, in order to offer best practice, it is imperative to ensure that a timely and proper diagnosis is made to confirm stroke and exclude SM to offer optimal therapy and aim to prevent irreversible brain damage.

FABS is a proposed scale used in discriminating SM from true strokes and consists of 6 variables (1 point for each) based on clinical assessment: absence of facial droop, negative history of atrial fibrillation, age < 50 years, systolic blood pressure < 150 mmHg on admission, history of seizures and isolated sensory deficits [9]. A score of ≥ 3 could help identify SM with a sensitivity of 90% and specificity of 91%. More recently, the simplified FABS (sFABS) demonstrated excellent discrimination and good calibration to predict SM in a Chinese stroke population receiving t-PA [10]. In Tanzania, there is paucity of data on the magnitude and outcomes of SM. We therefore aimed at investigating the proportion of SM among patients clinically diagnosed with stroke, assess the usefulness of the sFABS scale for predicting SM and determining the short-term outcomes of SM in a tertiary teaching hospital in Dar es Salaam, Tanzania.

Methods

Study design and population

This prospective study was conducted at Muhimbili University of Health and Allied Science Academic Medical Center (MAMC), a tertiary public teaching hospital in Dar es Salaam, Tanzania. MAMC offers super specialized medical care to all specialties, in the capital city of Dar es Salaam and receives referred patients from both public and private hospitals from all over the country.

All study patients were enrolled from June 2018 to January 2019. Those aged ≥18 years with a WHO clinical criteria for stroke, defined as the rapidly developing clinical signs of focal or global disturbance in cerebral function lasting more than 24 h or leading to death with no apparent cause other than that of vascular origin [1] were consecutively recruited and enrolled after obtaining written informed consent. Case ascertainment was accurate as all patients presenting to the emergency department with suspected stroke are transferred directly to the medical ward. An interviewer based structured questionnaire was administered to the patients or their caregivers in the event the patient was unable to communicate. The questionnaire captured socio-demographic information, mobile numbers and past medical/drug history for hypertension, diabetes mellitus, cardiac disease. We also inquired for smoking and alcohol consumption. The date of onset of stroke symptoms and date of arrival to the hospital was recorded in the questionnaire. Physical assessment of the patients was performed and included a focused neurological examination, measurement of waist-hip circumference and blood pressure (BP) on admission. Waist-hip ratio was computed and interpreted according to the WHO guidelines: in males the ratio of ≥0.90 and females ≥0.85 was regarded as substantially increased [11]. Subsequent BP readings were spaced 5 min apart and an average blood pressure was computed. The sFABS scale consisting of 4 items with 1 point in each item present: absence of facial droop, negative history of atrial fibrillation, age < 50 years and systolic blood pressure < 150 mmHg at presentation, was used to predict SM. Similar to a previous study [10], a score of ≥3 was tested for accuracy in the diagnosis of SM. Venous blood was collected aseptically from the cubital fossae and analysis was done for blood glucose levels, chemistry profile and complete blood count. Lumbar puncture was performed in patients with suspected meningoencephalitis. Brain imaging was used to confirm the diagnosis of stroke and these were termed as true strokes with either a non-contrast brain computed tomography (CT) scan or Magnetic Resonance Image (MRI) interpreted by a senior Radiologist. Contrast was administered in those with neuroimaging findings...
suspicious of tumor. A 12-lead electrocardiogram (ECG) using Bionet machine was performed on the patients to look for evidence of atrial fibrillation. Stroke severity was assessed using the National Institute of Health Stroke Scale (NIHSS) on the day of admission and the Modified Rankin Scale (mRS) was used to assess for disability at 24 h, 72 h, 7 days, 14 days and 30-days post-stroke [1]. Patients were contacted by phone if they had been discharged by these time points. The study was conducted and reported using strengthening the reporting or observational studies in epidemiology (STROBE) methodology.

Data analysis
Data was transferred from the questionnaires and entered into SPSS version 20.0 for analysis. Continuous variables were summarized and presented as means and standard deviation (SD) or medians with Interquartile Range (IQR). Comparisons of clinical presentation and 30-day outcomes among SM and true strokes were summarized as proportions and computed using Pearson’s Chi square test or Fisher’s exact test. The sFABS scale for prediction of stroke mimics was also analyzed for the purpose of determining its usefulness in the Tanzanian population. A $p$ value of $<0.05$ was used to denote statistical significance.

**Results**
Between June 2018 to January 2019, there were a total of 1403 admissions in the general medical ward, 729 (52%) females, out of which 430 (30.6%) patients met the WHO clinical criteria for stroke. We excluded 67 (16%) of patients for the following reasons: unable to consent, experienced recurrent strokes and those who did not complete brain imaging. We recruited the remaining 363 (92.4%) with a clinical diagnosis of first ever stroke and completed neuroimaging. The proportion of SM and true strokes are summarized in Fig. 1. The distribution of stroke subtype among patients with true strokes is summarized in Fig. 2: ischemic stroke accounted for 224 (66%), 109 (32%) had hemorrhagic stroke and 6 (2%) had mixed lesions (both ischemia and hemorrhage).
Demographic characteristics

Demographic characteristics of the SM and true strokes are summarized in Table 1. The mean age for SM and true strokes was 65.8 ± 15 and 57.9 ± 16 years, \( p = 0.024 \) respectively. The mean systolic and diastolic blood pressures were 126 ± 15 and 155.6 ± 21 mmHg, \( p < 0.001 \) and 77.2 ± 14 and 95.6 ± 11 mmHg, \( p < 0.001 \) for SM and true strokes respectively. The mean NIHSS score for SM and true strokes was 21.7 ± 8 and 21.2 ± 9, \( p = 0.770 \). SM were less likely to have premorbid hypertension 7 (29.2%) vs 263 (77.6%), \( p < 0.001 \) and increased waist-hip ratio 9 (37.5%) vs 270 (79.6%) \( p < 0.001 \) in SM and true strokes respectively.

### Table 1 A comparison of demographic characteristics between stroke mimics and true strokes

|                          | Stroke Mimics  | True strokes | Total       | \( P \) value |
|--------------------------|----------------|--------------|-------------|---------------|
| Age Mean ± SD            | 65.8 ± 15      | 57.9 ± 16    | 58.2 ± 16   | 0.024         |
| Female                   | 15 (62.5)      | 191 (56.3)   | 206 (56.7)  | 0.556         |
| Ever married             | 24 (100)       | 309 (91.2)   | 333 (91.7)  | 0.242         |
| Resident of Dar es Salaam| 12 (50)        | 265 (78.2)   | 277 (76.3)  | 0.002         |
| Insurance                | 5 (20.8)       | 92 (27.1)    | 97 (26.7)   | 0.5           |
| Medical History:         |                |              |             |               |
| Previous Hypertension    | 7 (29.2)       | 263 (77.6)   | 270 (74.4)  | < 0.001       |
| Previous Diabetes        | 1 (4.2)        | 61 (18)      | 62 (17.1)   | 0.095         |
| Previous Cardiac disease | 1 (4.2)        | 15 (4.4)     | 16 (4.4)    | 0.714         |
| Ever smoked              | 1 (4.2)        | 22 (6.5)     | 23 (6.3)    | 0.540         |
| Ever consumed alcohol    | 5 (20.8)       | 68 (20.1)    | 73 (20.1)   | 0.927         |
| Increased waist-hip ratio| 9 (37.5)       | 270 (79.6)   | 279 (76.9)  | < 0.001       |
| Atrial Fibrillation      | 2 (8.3)        | 20 (5.9)     | 22 (7.4)    | 0.695         |
| Systolic blood pressure  | 126 ± 15       | 155.6 ± 21   | 153.7 ± 22  | < 0.001       |
| Diastolic blood pressure | 77.2 ± 14      | 95.6 ± 10    | 94.3 ± 12   | < 0.001       |
| NIHSS Mean ± SD          | 21.7 ± 8       | 21.2 ± 9     | 21.2 ± 8    | 0.770         |
| Arrival to hospital from symptom onset: | | | | |
| Within 1 day             | 12 (50)        | 134 (39.5)   | 158 (40.2)  |               |
| Day 2 to 6               | 11 (45.8)      | 168 (49.6)   | 194 (49.4)  | 0.438         |
| > 7 days                 | 1 (4.2)        | 37 (10.9)    | 38 (10.5)   |               |

SD Standard deviation, NIHSS National Institute of Health Stroke Scale
A comparison of neurological signs among patients with stroke mimics and true strokes

During neurological examination, SM were less likely to have cortical signs: extinction 4 (16.7%) vs 128 (37.8%), \( p = 0.038 \), hemianopia 5 (20.8%) vs 153 (45.1%), \( p = 0.020 \) and supra-nuclear facial paralysis 6 (25%) vs 210 (61.9%), \( p < 0.001 \) for SM and true strokes respectively as shown in Table 2.

The use of sFABS scale among patients with stroke mimics and true strokes

Table 3 summarizes the distribution of sFABS score among patients with SM and true strokes. sFABS score of \( \geq 3 \) could identify patients with SM with a sensitivity and specificity of 38 and 80% respectively, while a sFABS score of \( \geq 2 \) had a sensitivity of 96% and specificity of 43%.

Causes of stroke mimics

The most frequent causes of SM were brain tumors 6 (25%), followed by meningoencephalitis 4 (16.7%), epileptic seizures 3 (12.5%) and brain atrophy 3 (12.5%) as summarized in Fig. 3.

Outcomes

The overall mortality at 30 days among the 363 patients was 209 (59.8%). SM had a shorter length of hospital stay of 6.2 ± 3.5 compared to 9.1 ± 16.1 days, \( p = 0.009 \) in true strokes. In survivors, there were no significant differences in functional independency or mortality at 30 days (summarized in Table 4). There were 15 (4.1%) lost to follow up following discharge; 2 (8.2%) SM and 13 (3.8%) true strokes. Patients lost to follow-up were excluded from the outcome analysis.

Discussion

The present study indicates that 6.6% of all admitted patients who initially met the WHO stroke definition for first event had SM as a final diagnosis. Our findings mirror those described by Forster A et al., where 6.4% of patients with suspected stroke had SM [12]. Other authors have reported higher proportions of SM (up to 30%); these variabilities in proportions could be attributed to the number of study patients enrolled, the setting and duration of the study, ours having fewer patients and a shorter study duration [3]. Nonetheless, our findings are alarming considering the majority of the countries in SSA (including Tanzania) have limited access to brain imaging. A one-year hospital based retrospective study conducted in three public tertiary hospitals in Zimbabwe included 450 patients with a clinical diagnosis of stroke. Of these, only 39.4% of the study patients underwent CT scans. They suggested that lack of resources and health insurance coverage could explain this low uptake of CT scans. Therefore, this signals the utmost importance of integrating such services to facilitate an early and accurate diagnosis, particularly when stroke presentations are associated with such high mortality (approximately 60% at 30 days in this study). Wardlaw et al., found that early stroke imaging with CT is the least costly strategy and associated with higher quality adjusted life years [14].

It is evident that the reliability of clinical history of an abrupt onset of focal or global neurological deficits such as hemiplegia, seen in more than half of the SM in this current study has its limitations. Some reports indicate that certain clinical features are highly suggestive of a SM and should be factored in when evaluating a suspected stroke patient [3, 15]. Such factors include the
paucity or absence of modifiable risk factors for stroke in clinical history [3]. In our study, the presence of pre-morbid hypertension and increased waist-hip ratio (makers of cardiovascular disease) were statistically less prevalent among SM compared to true strokes. Further, mimics were more likely to have a normal systolic and diastolic BP on admission compared to true strokes. Finally, cortical signs such as extinction, hemianopia and supra-nuclear facial paralysis were less-suggestive of SM. Our findings are similar to what was observed by Okano et al. and Yahia et al., were a systolic blood pressure < 140 mmHg on admission was independently associated with having a SM and cortical signs were described as pathognomonic for true strokes respectively [15, 16].

In this study, the use of sFABS score ≥ 3 in predicting SM among patients with suspected strokes, showed a low sensitivity (38%) and moderate specificity (80%). This contrasts with a sensitivity and specificity of 86 and 92% respectively when used in a Chinese population [10]. The low sensitivity of the scale observed in the current study could be explained by the fact that the sFABS scale was developed among those undergoing thrombolysis and therefore the current study population differs. Likewise, the demographics of the patients could be another factor; in the current study SM were elderly (mean age of 65 ± 15 years) and were therefore more likely to have lower scores. A sFABS score of ≤ 2 was observed in 62.4%, accounting for this low sensitivity. Our results are contrary to previous studies, were SM occur more commonly at younger ages and are less prevalent in the older population [16, 17]. Even after adjusting the cut off score to ≥ 2 in predicting SM, the sensitivity observed went as high as 96% with a low specificity of 46% making it unsuitable for use in clinical practice in our setting. Similarly, the biracial differences could also contribute to these discrepancies. We recommend further studies that can help develop clinical scales used for predicting SM feasible in SSA. This may help with better resource utilization, by identifying those most likely to have SM and thus benefit from imaging and other tests.

Our findings are in line with other authors, who have described brain tumors (observed in 25%) and epileptic seizures (12.5%), as the most common causes of SM [7, 15]. However, the high proportion of meningoencephalitis (16.7%) observed could be explained by the aging process which is associated with a greater propensity to contracting infections either due to physiological immunosenescence or having an underlying chronic disease.

Mortality was high at 30 days in our study, occurring in more than 50% of the patients with mimics, compared to mortality of 8.8% reported by Okano et al. [15]. The high mortality in our study could be explained by the fact that stroke mimics presented with severe disease on admission, with a mean NIHSS score of 21.7 ± 7.9 compared to a mean NIHSS score of 7.3 ± 8.9 observed in the former study [15].

**Table 4** A comparison of outcomes among patients with stroke mimics and true strokes

| Outcome                                | Stroke mimics        | True strokes      | Total          | P value |
|----------------------------------------|----------------------|------------------|----------------|---------|
| Length of hospital stay                | 6.2 ± 3.5            | 9.1 ± 16.1       | 8.9 ± 15.9     | 0.009   |
| Died at 30 days                        | 12 (54.5)            | 196 (60.1)       | 208 (59.8)     | 0.295   |
| Independency at 1 month (amRS 0–2)    | 1 (10)               | 19 (14.6)        | 20 (14.3)      | 0.522   |

*amRS Modified Rankin Scale*
Our study had the following limitations; it was a single center with a small sample size and a short follow up period therefore results cannot be generalized.

Conclusions
In the present study the proportion of SM among patients clinically diagnosed with stroke was 6.6% and brain tumors was a common etiology. SM were less likely to have cardiovascular risk factors and cortical signs during evaluation. We thus recommend further studies that can help develop clinical scales used for predicting SM in an African population.

Abbreviations
BP: Blood pressure; CT: Computed tomography; ECG: Electrocardiogram; FABS: Stroke mimic scoring system; IQR: Interquartile Range; MAMC: Muhimbili University of Health and Allied Science Academic Medical Center; MRI: Magnetic Resonance Imaging; mRS: Modified Rankin Scale; NIHSS: National Institute of Health Stroke Scale; SD: Standard deviation; SM: Stroke mimics; SSA: Sub Saharan Africa; STROBE: Strengthening the reporting of observational studies in epidemiology; t-PA: Tissue plasminogen activator; WHO: World Health Organization

Acknowledgements
Our gratitude goes to the study patients, Dr. Candida Moshiro for her statistical advice, Dr. Teddy Msielle and Mr. Akili Mawazo for analyzing the samples and finally Dr. Freddy Lyimo a senior radiologist for imaging interpretation.

Authors’ contributions
SSM and PM conceptualized and designed the study. SSM and KK collected the initial manuscript. All authors contributed to the draft and approved the final manuscript.

Funding
This study was funded by Catholic University of Health and Allied Sciences. The funder had no role in the design, analysis, final write up of the manuscript and decision to submit the paper for publication.

Availability of data and materials
The dataset used for analysis in this study is available from the corresponding author on reasonable request.

Ethics approval and consent to participate
Ethical clearance was obtained from Muhimbili University of Health and Allied Science Institutional review board approval number DA.287/298/01A/. Written informed consent to take part in the study was obtained from all patients or their next of kin prior to study enrollment. All patients were offered standard medical care following diagnosis.

Consent for publication
All patients or their next of Kin gave consent for publication for any clinical information or investigations collected that was requested for during the written informed consent process.

Competing interests
The authors declare that they have no competing interests.

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Received: 17 April 2020 Accepted: 1 July 2020
Published online: 07 July 2020

References
1. WHO Noncommunicable Diseases and Mental Health. The WHO STEPS approach to stroke surveillance report. 2005. Available from: https://www.who.int/nmh_surveillance/en/steps_stroke_manual_v1.2.pdf.
2. Nitka M. Stroke mimics. Emerg Med Clin North Am. 2012;30(3):795–804.
3. Menino JG, Luby M, Benson RT, Davis LA, Hsia AW, Latour LL, et al. Predictors of acute stroke mimics in 8187 patients referred to a stroke service. J Stroke Cerebrovasc. Dis. 2013;22(8):e397–403.
4. Zinkstok SM, Engelter ST, Genetichke L, Lyer PA, Ringleb PA, Arto Y, et al. Safety of thrombolysis in stroke mimics: results from a multicenter cohort study. Stroke. 2013;44(4):1080–4.
5. Onwuekwe JO, Ezeala-Adikaike BA, Ohaegbulam SC, Chikani MC, Amuta J, Uloh HN. Stroke mimics - study of CT images in Nigerian African stroke patients. J Neurol Sci. 2008;25(3):143–9.
6. Yew KS, Medicine F, Cheng VEM, Angeles L. Diagnosis of acute stroke. Am Fam Physician. 2015;91(8):528–36.
7. Communication S. Stroke mimics in patients with clinical signs of stroke. Cspian J Intern Med. 2017;8(3):213–6.
8. Walker RW, Viney R, Green L, Mawanswila M, Maro VP, Gjertsen C, et al. Trends in stroke admissions to a Tanzanian hospital over four decades: a retrospective audit. Trop Med Int Heal. 2015;20(1):1290–6.
9. Goyal N, Tsivgoulis G, Male S, Metter EJ, Iftikhar S, Kerro A, et al. FABS: an intuitive tool for screening of stroke mimics in the emergency department. Stroke. 2016;47(9):2216–20.
10. Qin X, Zhao S, Yin L, Dou H, Fu J, Wang Y, et al. Validation of simplified FABS scale to predict stroke mimics in a Chinese population undergoing intravenous thrombolysis. Clin Neurol Neurosurg. 2017;161:1–5.
11. World Health Organization. Waist circumference and waist-hip ratio: report of a WHO expert consultation. Geneva WHO; 2011. p. 1–39. (8–11 December 2008).
12. Förster A, Greibe M, Wolf ME, Szabo K, Hennerici MG, Kern R. How to identify stroke mimics in patients eligible for intravenous thrombolysis? J Neurol. 2012;259(7):1347–53.
13. Kaseke F, Stewart A, Gwanzura L, Hakim J, Chikwasha V. Clinical characteristics and outcomes of patients with stroke admitted in tertiary service. J Stroke Cerebrovasc Dis. 2013;22(8):e397–403.
14. Wardlaw J, Keir S, Seymour J, Lewis S, Sandercock P, Dennis M, et al. What is the best imaging strategy for acute stroke? Heal Technol Assess. 2004(8):1–100.
15. Okano Y, Ishimatsu K, Kato Y, Yamaga J, Kuvahara K, Okumoto K, et al. Clinical features of stroke mimics in the emergency department. Acute Med. 2016;5(3):241–8.
16. Yahia MM, Bashir S. Clinical characteristics of stroke mimics presenting to a stroke center within the therapeutic window of thrombolysis. Brain Neurorehabil. 2018;1(1):9.
17. Campello AR, Steinhauser EG, González AG, Martínez MS, Cerajería POC, Torres AZ, De, et al. Description of stroke mimics after complete intravenous thrombolysis. Neurologia (English Ed) 2019;34(1):7–13. doi: https://doi.org/10.1016/j.nrleng.2016.10.004.

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