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Nadeem Kassam
Omar Aziz
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Hanifa Mbithe

See next page for additional authors
Authors
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Nadeem Kassam1
Omar Aziz1
Eric Aghan2
Samina Somji1
Hanifa Mbithe1
Mustafa Bapumia3
Robert Mvungi3
Salim Surani4

1Department of Internal Medicine, Aga Khan University Medical College East Africa (Dar), Dar es Salaam, Tanzania;
2Department of Family Medicine, Aga Khan University Medical College East Africa (Dar), Dar es Salaam, Tanzania;
3Department of Internal Medicine and Cardiology, Aga Khan Hospital, Dar es Salaam, Tanzania; 4Department of Medicine & Pharmacy, Texas A & M University, College Station, TX, USA

Abstract: Smartwatches like the Apple Watch have been on the rise worldwide and their use is gaining popularity in developing countries. Their ability to detect dysrhythmias is well documented. Present practice discourages the use of these devices as a diagnostic tool. Nevertheless, atypical findings from these devices should be clinically investigated. This case demonstrates an eventual diagnosis of supraventricular tachycardia (SVT) based on an Apple watch alert which was subsequently confirmed by electrophysiological evaluation.

Keywords: supraventricular tachycardia, Apple watch, Tanzania, electrophysiology study

Introduction

Global sales of wearable smartwatches have grown at an annual rate beyond 20%, with approximately half a billion active users in 2020 which is expected to double by 2022.1 The incorporation of photoplethysmographic (PPG) sensors into smartwatches has enhanced extensive community-based heart rate monitoring. The latest version of smartwatches incorporate the PPG sensor that can identify pulse rate and regularity offering precise results directly to the user. Despite having a paucity of scientific evidence to support the use of these devices to detect either symptomatic or malignant arrhythmias. There are numerous anecdotal reports of patients saved by their smartwatches which were able to detect and warn the user of an arrhythmia.1 We hereby present a case of a 42-year-old African male who was alerted to a possible arrhythmia by his smart watch and was subsequently diagnosed to have supraventricular tachycardia (SVT).

Case Report

A 42-year-old male with well-controlled hypertension on once-daily Losartan 50 mg presented with acute chest discomfort associated with palpitations and lightheadedness. He was at home watching television when these symptoms started acutely and continued for almost 10 minutes. In addition, his Apple Watch (Apple Watch Series 4) recorded a high resting heart rate of 210 beats/min (Figure 1). Alarmed at the elevated heart rate, he hurried to the emergency department. On examination, the patient was conscious, awake and oriented with warm peripheral extremities. Initial vitals at the emergency department revealed tachycardia of 184 beats/minute, blood pressure of 120/80 mmHg and respiratory rate of 16 cycles per minute; his oxygen saturation was 99% on room air. Systemic examination was unremarkable. A 12 lead electrocardiogram (ECG) was performed (Figure 2) and revealed supraventricular tachycardia at the rate of 184 b/min. The patient was
given intravenous Adenosine 6 mg and then two additional doses of 12 mg via cubital peripheral line but failed to revert to a normal sinus rhythm. At this point, he developed hypotension and was quickly electrically cardioverted (synchronized DC 50 Joules) and successfully reverted to normal sinus rhythm as seen in the repeat ECG below (Figure 3). Subsequent workup with a complete blood count, thyroid panel, repeated cardiac enzymes, Troponin I, electrolytes, renal functions were all normal. 2D echocardiogram revealed normal left ventricular systolic (ejection fraction 60%) and diastolic functions with normal-sized chambers. Coronary angiography demonstrated no evidence of epicardial coronary artery disease. The patient was discharged on daily Nebivolol 5 mg, Atorvastatin 20 mg and continued his daily Losartan 50 mg with a plan of electrophysiological studies (EPS) to establish the possible cause of the symptomatic supraventricular tachycardia (SVT). With no EPS studies available in our country, he subsequently flew to India after four weeks and underwent an electrophysiology evaluation which revealed inducible tachycardia on pacing protocols. In addition, electrophysiological mapping showed reproducible SVT, which was orthodromic atrio-ventricular reentrant tachycardia (oAVRT) utilizing parahisian accessory pathway as retrograde conduction. A ten second ablation was done on the non-coronary

Figure 1 Apple watch record of the patient's heart rate.

Figure 2 EKG showing supraventricular tachycardia.
sinus (NCS), which was considered the best option but had no effect on the tachycardia. A test ablation was considered on the parahisian pathway but was deemed to be associated with too high of a risk for permanent pacing, which the patient declined. The patient was discharged on medical management with the option of cryoablation for the failure of medical therapy. A 6 and 12 month’s follow-up has been uneventful.

Discussion
To our knowledge, this is the first published case report in Sub-Saharan Africa whereby the Apple Watch was an integral part of the diagnostic process. Many patients who present to health care facilities reporting awareness of heartbeat require extensive workup: including ambulatory rhythm monitoring, noninvasive imaging, pharmacotherapy and invasive electrophysiological (EP) procedures to reach diagnosis. Literature from the African context shows that even though patients may present for care, most facilities are ill-equipped and lack necessary diagnostic capabilities both in human resources and equipment. This advancement in technology has shed light on the medical world and can therefore be considered a significant adjunct to conventional rhythm monitoring. Nevertheless, it is a dangerous practice to solely depend on it to detect heart conduction abnormalities. This technology has led to an extensive assessment of smartwatches to identify atrial fibrillation. The findings of this study were clinically relevant since 34% of those who received an irregular pulse notification were subsequently diagnosed to have atrial fibrillation on ECG patch readings. Additionally, case series have also been published whereby two patients’ actual watch-based 3-lead ECG tracings corresponded to the traditional ECGs demonstrating ST-elevation myocardial infarction (STEMI), confirming the potential ability of this device also to detect myocardial ischemia. Only one validation study has assessed the accuracy of the wrist-worn wearable device on 51 patients who were scheduled for EPS due to a history of paroxysmal supraventricular tachyarrhythmia (SVT) or paroxysmal palpitations. This study showed an acceptable accuracy of these devices to measure baseline and induced SVT.

Conclusion
This case highlights the importance of wearable gadgets to detect asymptomatic arrhythmia. Atrial fibrillation is
considered most prevalent among hypertensive patients in Tanzania. The use of such devices will therefore aid in seeking prompt care and management. However, false-positive notifications can trigger anxiety and increase unnecessary hospital visits. The American and European heart rhythm society has not approved this technology as a diagnostic tool rather a nidus for the community to seek medical advice when notified.

Declaration of Patient Consent
The authors certify that patient consent was obtained. The patient has given his consent for his images and other clinical information to be reported in the journal. The patient understands that his name and initials will not be published. The obtained consent was also forwarded to the institutional ethical committee. Institutional approval was not required to publish the case details.

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Disclosure
The authors report no conflicts of interest for this work.

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