Cardiac Arrhythmias in Africa
Epidemiology, Management Challenges, and Perspectives

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ABSTRACT

Africa is experiencing an increasing burden of cardiac arrhythmias. Unfortunately, the expanding need for appropriate care remains largely unmet because of inadequate funding, shortage of essential medical expertise, and the high cost of diagnostic equipment and treatment modalities. Thus, patients receive suboptimal care. A total of 5 of 34 countries (15%) in Sub-Saharan Africa (SSA) lack a single trained cardiologist to provide basic cardiac care. One-third of the SSA countries do not have a single pacemaker center, and more than one-half do not have a coronary catheterization laboratory. Only South Africa and several North African countries provide complete services for cardiac arrhythmias, leaving more than hundreds of millions of people in SSA without access to arrhythmia care considered standard in other parts of the world. Key strategies to improve arrhythmia care in Africa include greater government health care funding, increased emphasis on personnel training through fellowship programs, and greater focus on preventive care. (J Am Coll Cardiol 2019;73:1009–10) © 2019 by the American College of Cardiology Foundation.

Any countries face growing, unmet health care needs, and pressures on health systems are expected to increase as a result of demographic, epidemiological, and health transitions (1). Health care delivery across the African continent is heterogeneous, with few countries providing universal health coverage. Among 55 countries in Africa, only 8 provide universal health coverage through a national health insurance program: Algeria, Botswana, Gabon, Ghana, Mauritius, Rwanda, Tunisia, and Zambia (2). In most countries, health care services are primarily paid out of pocket by household incomes rather than by public or commercial health insurance programs (3). In 2014, between 40% and 60% of total health expenditures were funded by households in one-half of Sub-Saharan Africa (SSA) countries (3). Although most governments have committed to increasing their public health spending to at least 15% of the government’s budget in line with the African Union’s 2001 Abuja Declaration (3), government health expenditures diminished in one-half of the African countries between 2002 and 2014 (1).

Cardiovascular diseases, including cardiac arrhythmias, are a major public health problem in low- and middle-income countries, to which almost all SSA countries belong. There is a huge disparity in cardiovascular standard of care between the developed countries and the low- and middle-income countries. Because of the high cost of diagnostic

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and therapeutic interventions in electrophysiology, cardiac arrhythmia may possibly be the most neglected area of cardiology. To address this situation, we review the current state of arrhythmia burden and care, identify critical needs, and suggest some potential solutions. We recommend a hierarchy of cooperative goals, which can be set across African countries, starting with education and care less dependent on high-cost items.

**Epidemiology**

**Atrial Fibrillation.** Atrial fibrillation (AF) is the most common arrhythmia worldwide, including Africa (4,5). From 1990 to 2013, deaths from AF increased 196% (4). According to some projections, there will be more people with AF in Africa than in either China, the United States, or India by 2050 (Figure 1).

Rheumatic heart disease and endomyocardial fibrosis are still prevalent in Africa, affecting mainly adolescents and young adults. They are associated with a high prevalence of AF (4–6). AF is found in >30% of the chronic cases of endomyocardial fibrosis (6). In the REMEDY (Global Rheumatic Heart Disease Registry), AF was present in 21.8% of patients with rheumatic heart disease (7), which is consistent with the findings of the Heart of Soweto study, which showed a 13% prevalence of AF among patients with valvular heart disease (8).

A study from urban Cameroon was the first attempt to describe the clinical characteristics, treatment, and prognosis of patients with AF in Africa (9). Of 172 patients with AF, 22.7% were paroxysmal, 21.5% were persistent, and 55.8% were permanent. In this 1-year follow-up study, patients experienced extremely high mortality and morbidity rates: 26 of 88 patients (30%) died; among the survivors, 16% had a nonfatal stroke and 26% developed decompensated heart failure (HF).

In African HF registries, 16% of acute HF patients in THESUS-HF (Sub-Saharan Africa Survey on Heart Failure) (10) and 18% of chronic HF patients in TaHeF (Tanzanian Heart Failure Study) (11) had AF. Both studies present a lower prevalence of AF compared with that reported in developed countries (5). The reason for this is not clear and may be related to improved survival and increased chronicity of HF in patients in the high-income countries or perhaps because the African patients were younger than their Western counterparts. In the international RE-LY (Randomized Evaluation of Long-Term Anti-coagulation Therapy) registry, patients presented with AF or atrial flutter to an emergency department.

African patients had about 3 times more hospital readmissions (34%) than others; their 1-year stroke rate was 8% compared with 2% in North America, Western Europe, and Australia; and the 1-year death rate was 20% compared with 10% on the other continents (12).

**Inheritable Arrhythmogenic Heart Diseases.** There is a paucity of data on inherited arrhythmogenic cardiac diseases due to the advanced workup required to establish the diagnosis. Hence, the published data is relatively limited regarding this topic (Table 1).

**Arrhythmogenic Right Ventricular Cardiomyopathy/Dysplasia.** A rare heart muscle disorder characterized by the presence of fibrofatty tissue and ventricular electrical vulnerability related to sudden death. In 2009, the largest cohort of 50 patients in Africa was reported by the Arrhythmogenic Right Ventricular Cardiomyopathy/Dysplasia Registry of South Africa (13,14). The disease was observed in all races and ethnic groups, patients usually became symptomatic by their third decade of life, and the 1- and 5-year mortality was 2.8% and 10%, respectively. The relatively high mortality was believed to be the result of underutilization of implantable cardioverter-defibrillators (ICDs) in the country (11). In Tunisia, 12 sudden cardiac death (SCD) patients with suspected arrhythmogenic right ventricular cardiomyopathy/dysplasia were studied (15). Autopsies revealed that 9 patients met the morphological criteria of the disease (adipocyte infiltration with fibrosis replacement) (15).

**Long QT Syndrome** is one of the most common cardiac channelopathies, characterized by prolonged ventricular repolarization and life-threatening ventricular arrhythmias called “torsades de pointe,” which may lead to syncope and cardiac arrest. The first published case of long QT syndrome on the continent came from South Africa in 1992 (16). The case was a 7-year-old child from Senegal who presented with recurrent episodes of exercise-associated syncope, a QTc of 627 ms, and a Schwartz score of 6. Neither syncope nor ventricular arrhythmia recurred on propranolol until discharge 2 weeks later. Unfortunately, no genetic testing was done. Despite such scant case reports, there is no substantial understanding of the prevalence of this condition in Africa.

**Brugada Syndrome** is an inherited arrhythmogenic disorder phenotypically manifested by syncope, malignant ventricular arrhythmias or SCD, and coved-type ST-segment elevation in the right precordial leads on the electrocardiogram (ECG). Although there
are few data on the prevalence of Brugada syndrome in the general population of Africa, Ouali et al. (17) reported 24 confirmed cases in Tunisia and found that the patients had a clinical profile like those from Western countries. Bonny et al. (18) reported on 5 black African patients with similar phenotypic/genotypic features as non-Africans.

A few sporadic cases of hypertrophic cardiomyopathy in Africans have been reported (19). To our knowledge, other conditions such as short QT syndrome, catecholaminergic polymorphic ventricular tachycardia, noncompaction cardiomyopathy, and a malignant form of early repolarization syndrome have not been reported in Africa. Clearly there is a need for improved recognition and reporting of the incidence of the inherited arrhythmia syndromes in Africa.

SCD is defined as death from an unexpected circulatory collapse, usually because of a cardiac arrhythmia, occurring within an hour of symptom onset. SCD data from most African countries are generally single-center retrospective registries rather than prospective population-based studies (Table 1). The first population-based SCD study in SSA (20) showed an incidence similar to China (21) and higher than in Ireland (22). The crude incidence rate in the Cameroonian population above age 18 years was 31.3/100,000 person-years, while the age-standardized rate across Africa was 33.6/100,000 person-years (20). Because of cultural misconceptions about unexpected deaths, which are seen as spiritual rather than medical events, post-mortem autopsies is rarely performed. Therefore, only speculations on the etiologies of SCD have been made. Inherited cardiac arrhythmias may be the cause of most of the SCD in areas where more than one-half of the population are age <25 years. In the Douala-SCD study, the number of deaths above age of 40 years was not greater than those under 40 years (20), suggesting that the proportion of patients with inherited arrhythmogenic diseases may be higher in the African populations compared with Western populations, in which ischemic heart disease (IHD) accounts for about 80% of all SCD in adults. Alternative conditions affecting such individuals under 40 years of age in Africa cannot be excluded.

The epidemiology of sports-related SCD is not well established in Africans. However, Allouche et al. (23) described 32 cases of SCD in athletes who underwent autopsies. The authors found a prevalence pattern similar to that reported in non-African populations: hypertrophic cardiomyopathy (41%) and arrhythmogenic right ventricular cardiomyopathy/dysplasia (13.6%) among those age <35 years, and ischemic heart disease (90%) in athletes age >35 years.

SECONDARY ARRHYTHMOGENIC HEART DISEASES. Ischemic heart disease. Historically, IHD has been considered uncommon in Africa (4). Despite the epidemiological transition, IHD continues to be responsible for a minority of SCDs in SSA (24–27). In North Africa, IHD was much more prevalent and accounted for almost 60% of 361 autopsied SCD fatalities in Tunisia (28).

Parasitic-related cardiac arrhythmias. Some parasitic infections may affect the myocardium, pericardium, and pulmonary vasculature and produce cardiac manifestations and rhythm disturbances. Human African Trypanosomiasis is caused by Trypanosoma brucei gambiense (in West and Central Africa, predominantly in the Democratic Republic of Congo) and Trypanosoma brucei rhodesiense (in East and South Africa, predominantly in Tanzania and Uganda). These flagellate parasites can cause myocarditis and prolongation of the PR, QRS, and QT intervals, with the associated risk of fatal ventricular arrhythmias. Cysticercosis, caused by Taenia solium, is a cestode parasite most commonly found in Southern Africa that can cause conduction abnormalities and trigger arrhythmias. Schistosomiasis (Schistosoma mansoni and S. Haematobium in West Africa) causes pulmonary hypertension, cardiac
arrhythmias, and SCD. The pathogenesis of parasitic-related arrhythmias is poorly understood, but inflammation, myocarditis, and cell death with fibrosis replacement have been postulated (29). The true impact of parasitic disease in the genesis of arrhythmias in Africa may be much higher than in other parts of the world, and merits further exploration as part of a more general registry on cardiac rhythm disturbances on the continent.

**DIAGNOSTIC AND THERAPEUTIC RESOURCES**

Accurate diagnosis and treatment of cardiac arrhythmias are facilitated by ambulatory rhythm monitoring, noninvasive imaging, pharmacotherapy, and invasive electrophysiological (EP) procedures. To ascertain how patients are managed across the continent, the Pan-African Society of Cardiology (PASCAR) Task Force on SCD conducted 2 online surveys from 2011 to 2017 (1,30). Cardiologists were invited to answer questions regarding the availability of diagnostic tools and treatments in their respective countries. Device manufacturers contributed data as well. Data from 33 of 55 African countries were analyzed and showed considerable heterogeneity of cardiac arrhythmia management across the continent (Central Illustration). A total of 5 countries (15%) did not have a single cardiologist, 10 (30%) had no pacemaker implantation services, and 18 (54.5%) had no electrophysiology laboratory (30). ECG, signal-averaged ECG, echocardiography, ambulatory ECG monitoring, exercise stress testing and head-up tilt-testing, and drugs are largely unavailable outside the major cities.

**PHARMACOTHERAPY.** As shown in the second PASCAR survey (30), intravenous anti-arrhythmic drugs are available only in a few teaching centers in the more developed countries, i.e., the North Africa region and South Africa. Digoxin is the most commonly used oral agent, mainly for AF rate-control and HF treatment. Rhythm-control strategy is rarely employed, with amiodarone used as the first line anti-arrhythmic medication.

**INVASIVE TREATMENTS.** Considerable heterogeneity in the access to invasive arrhythmia care was observed across Africa (Table 2). About one-third of African countries do not perform pacemaker implantations, leaving hundreds of millions of people without adequate access to the treatment of brady-cardia and especially heart block (30).

In 2014, the median pacemaker implantation rate in Africa was 2.66 per million population per country, which is 200-fold lower than in Europe (1). Extreme variation in medical charges (up to 1,000-fold) was observed across countries with an inverse correlation between implantation rates and the procedural fees standardized to the gross domestic product per capita (1).

The lack of national registries on cardiac implantable electronic devices (CIEDs) makes it difficult to evaluate the impact of ICD implantation on primary prevention of SCD, or the role that cardiac resynchronization therapy (CRT) plays in HF management in Africa. HF is a major cause of hospitalization in Africa. HF mortality rate is high due to inadequate treatment (9–31). Indeed, as reported in

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**TABLE 1** Nonexhaustive List of Published Data on Sudden Cardiac Death in Africa

<table>
<thead>
<tr>
<th>Main Etiology</th>
<th>First Author (Ref. #)</th>
<th>Title</th>
<th>Country</th>
<th>Journal, Year</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unknown</td>
<td>Bonny, et al. (20)</td>
<td>Epidemiology of sudden cardiac death in Cameroon</td>
<td>Cameroon</td>
<td>Int J of Epidemiology, 2017</td>
</tr>
<tr>
<td>IHD</td>
<td>Rotimi, et al.</td>
<td>Sudden unexpected death from cardiac causes in Nigerians: a review of 50 autopsied cases</td>
<td>Nigeria</td>
<td>Int J Cardiol, 1998</td>
</tr>
<tr>
<td>Hypertension</td>
<td>Akinwusi, et al.</td>
<td>Pattern of sudden death at Ladoke Akintola University of Technology Teaching Hospital, Osogbo, South West Nigeria</td>
<td>Nigeria</td>
<td>Vasc Health Risk Manag, 2013</td>
</tr>
<tr>
<td>IHD, PPCM, DCM</td>
<td>Talle, et al.</td>
<td>Sudden Cardiac Death: Clinical Perspectives from the University of Maiduguri Teaching Hospital, Nigeria</td>
<td>Nigeria</td>
<td>World J Cardiovasc Dis, 2015</td>
</tr>
<tr>
<td>HCM</td>
<td>El-Saiedi, et al.</td>
<td>Hypertrophic cardiomyopathy: prognostic factors and survival analysis in 128 Egyptian patients</td>
<td>Egypt</td>
<td>Cardiol Young, 2014</td>
</tr>
</tbody>
</table>

CMP = cardiomyopathy; DCM = dilated cardiomyopathy; HCM = hypertrophic cardiomyopathy; IHD = ischemic heart diseases; LQTS = long QT syndrome; PPCM = postpartum cardiomyopathy; RDC = République démocratique du Congo.
the PASCAR surveys (1,30), the implantation rate of ICDs is extremely low in all countries where this service is available, accounting for 12.8 and 11.7 devices per million population in Tunisia and South Africa, respectively. To the best of our knowledge, there are no published data on the indications for nonischemic ICD implantations in Africa, even though Cabral et al. (19) reported the first ICD placement in Central Africa in a hypertrophic cardiomyopathy patient in 2016. However, in the Euro-survey CRT 2, data from Algeria corroborates the trend reported in the first PASCAR survey (1) showing that CRT-P were more frequently implanted than CRT-D.

As shown in Figure 2, EP studies including complex catheter ablations are rarely available in Africa except in South Africa and the North African countries (1). Cryoablation was available in a few countries (i.e., Algeria, Egypt, Morocco, Namibia, South Africa, and Tunisia).

REASONS BEHIND INADEQUATE STATE OF CARDIAC ARRYTHMIA MANAGEMENT

SOCIOECONOMIC CONDITIONS. Comprehensive management of cardiac arrhythmias in African countries is limited by the following challenges:

1. Lack of political will by many governments of the region to establish and support hospitals, train the workforce, and purchase and maintain equipment necessary to provide comprehensive EP services.
Lack of facilities and a low number of trained practitioners to detect and manage cardiac arrhythmias is one of the main problems with EP services in Africa. The PASCAR surveys (1,30) highlighted the deficiencies in the human resources and facilities for arrhythmia. The number of invasive cardiac physicians and technicians per population is quite low. Besides Mauritius and Tunisia, the number of pacemaker implantation centers is ≤1 per million population. In one-half of the countries, there were no CRT implants. Tunisia had the highest rate of arrhythmia center density with 0.81 centers per million population. Hence, millions of people live without access to cardiac arrhythmia treatment.

POOR ACCESS TO ANTIARRHYTHMIC DRUGS AND ANTICOAGULANTS. In the urban Cameroon study (8), rate control was the principal strategy for treatment of AF, chosen in 84% of patients, most of which...
were treated with digitalis; the remaining 16% were treated by a rhythm-control approach with amiodarone or sotalol. Oral anticoagulants were prescribed in only 34% of eligible patients, despite high CHADS2 scores and a high rate of rheumatic heart disease. This scenario is common on the continent. Several barriers limit appropriate management of AF in Africa: 1) unavailability of most antiarrhythmic drugs on the African market; 2) the relatively high cost of vitamin K antagonist follow-up using the international normalized ratio; 3) physician ambivalence toward prescribing vitamin K antagonist secondary to the perception of poor reliability of international normalized ratio results produced by the local laboratories; and 4) the high cost of non-vitamin K oral anticoagulants, and their lack of availability in many SSA countries (30,33). Educating general practitioners (who are the ones that manage cardiac arrhythmias in poor-resource areas) and even cardiologists to become more familiar with antiarrhythmic drugs, anticoagulants, and rate-controlling agents other than digitalis should be a priority. The availability and appropriate use of external electrical cardioversion or defibrillation are suboptimal (Table 2).

**INADEQUATE RESPONSE TO CARDIAC ARREST.** Out-of-hospital cardiac arrest is the most common presentation of sudden cardiac arrest according to the Douala SCD-study (20) and a Tunisian study (28), and it occurs mainly in the presence of a witness. However, cardiopulmonary resuscitation is rarely attempted, leading to the high rate of potentially preventable deaths (20,34). Reasons for the low cardiopulmonary resuscitation rate include: 1) a lack of basic knowledge about cardiopulmonary resuscitation in the general population; 2) a lack of...
appreciation for the urgency of the situation; 3) a lack of external defibrillators in public places; 4) a lack of minimal resuscitation equipment and emergency drugs (e.g., parenteral antiarhythmic drugs, atropine, and adrenaline) in ambulances; and 5) difficulties in reaching emergency medical service operators, partly because of an absence of toll-free emergency telephone services along with poor identification of the victim’s physical address owing to lack of uniformly agreed upon administrative city plans. We call for the standardization of cardiopulmonary resuscitation training for medical staff and the general public in schools and government offices to address the growing incidence of sudden cardiac arrest.

**NEEDS ASSESSMENT AND OPPORTUNITIES**

**BASELINE CARDIAC WORKUP.** There is an enormous need to improve the availability of basic diagnostic tools (electrocardiograms, ambulatory heart monitors, and echocardiograms), as well as for training general physicians and nurses to recognize common cardiac arrhythmias such as AF and to manage their complications. The use of vitamin K antagonist (and international normalized ratio monitoring), non-vitamin K oral anticoagulants, beta-blockers, angiotensin-converting enzyme inhibitors, and angiotensin receptor blocker agents should be standardized and promoted through educational symposia (35).

**CONNECTED CARDIOLOGY.** The African cardiac community should work with other cardiac societies to develop an internet-based network to connect cardiologists and general practitioners in Africa and to also connect with experts outside the continent. A system of electronically accessible electrocardiography (e-ECG) might help to detect actionable arrhythmias. PASCAR should also develop and recommend simple strategies and algorithms for the management of common arrhythmias, taking into consideration the peculiarities of the African health-care system (2-4,33).

**PEDIATRIC CARDIOLOGY.** There is increasing realization that the lack of facilities for sustainable pediatric cardiac arrhythmia services in the developing world results in significant and potentially treatable morbidity and preventable deaths (36). Particularly in SSA, the high burden of rheumatic heart disease- and endomyocardial fibrosis-related AF and unexplained premature infant deaths threaten to worsen the life-expectancy rate of this young African population. While pediatric patients with congenital heart disease undergo surgery in few countries of all subregions of the Continent, to the best of our knowledge, pediatric cardiac arrhythmia invasive procedures are available only in Algeria, Egypt, and South Africa. Children with repaired congenital heart disease-related arrhythmias do not have any access to specialized treatments elsewhere on the continent.

**FELLOWSHIP INITIATIVES.** PASCAR initiated a fellowship training program in cardiac pacing to provide the required expertise in SSA. The long-term objective of this program is to train a team for every country without a pacing service by 2030. Since the program was launched in 2015, 2 cardiologists from Sierra Leone and Kenya along with their nurses have been trained through a 6-month fellowship in Cardiac Pacing in Cape Town, South Africa. However, the program faces funding challenges. To attract funds and build additional capacity, PASCAR engaged in collaboration with the Heart Rhythm Society (Online Appendix) and CIED manufacturers.

Establishment of the Pan-African arrhythmia and pacing fellowship, endorsed by PASCAR and select African university medical centers, would provide physicians in Africa the opportunity to train locally at a lower cost. Further collaborations with the Asia-Pacific Heart Rhythm Society and European Heart Rhythm Association should also be encouraged. Collaborations with these professional groups should include donations of medical equipment as replacements along with appropriately recycled implantable devices (37).

**USE OF RECONDITIONED CIEDs.** CIEDs are one of the most cost-intensive treatments in the arrhythmia management armamentarium. CIED reuse has been proposed as a means of addressing the large disparity in access to this therapy. PASCAR has created a task force for pacemaker and ICD reuse. Along with the U.K.-based charity Pace 4 Life and the University of Michigan program My Heart Your Heart, PASCAR has worked to bring reconditioned CIEDs for patients in Africa. All reconditioned devices will be entered into a registry, and clinical outcomes will be reported periodically.

**UNIVERSAL HEALTH CARE.** Comprehensive universal health coverage and the development of health insurance policies should include coverage for arrhythmia care and SCD. Public-private partnership can be explored especially in bridging the existing gaps regarding facilities. Governments could develop policies to provide subsidies to make CIEDs available to patients who are at high risk of sudden death.

**MEDICAL EVACUATIONS.** A glaring example of misappropriation of resources and inequity in health care is medical evacuations. A concept unfamiliar to health professionals in high-income countries, many
African governments approve medical evacuations (i.e., transfers to tertiary health facilities abroad) for a small privileged segment of the society. This unseemly practice benefits the wealthy and government elites, is highly cost-ineffective, and misappropriates limited local resources that cannot be used for maximum benefit of very ill patients. As a health policy, such arbitrary allocation is unethical.

**STUDY LIMITATIONS**

The epidemiology of arrhythmia and its treatment has been understudied in Africa. There is lack of registries and cross-sectional and cohort studies. The published data available is of limited value, owing to its observational, single-center, or case report character. There is a great need for pan-African registries on cardiac rhythm disturbance management.

**CONCLUSIONS**

Public health strategies in many African countries do not adequately address the growing burden of cardiac arrhythmias. Millions of patients are left without access to treatments considered standard in high-income countries. The areas of focus should be: 1) increased health care financing and research for Africa-specific needs; 2) observational studies and registries to ascertain the disease patterns specific to the continent; 3) training of nurses, general practitioners, and cardiologists; and 4) collaboration among PASCAR cardiologists and other cardiologists and heart rhythm societies. Progress toward universal health coverage would further improve access to care and ensure equitable health improvement in the population.

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**KEY WORDS** Africa, cardiac arrhythmias, cardiovascular diseases, epidemiology, management

**APPENDIX** For the Heart Rhythm Society meeting minutes, please see the online version of this paper.