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Aziza Razki¹,², Eva Hong¹, Khalid Zerouali³, Houria Belabbes², Khadija Aitmouss², Aude Terrade¹, Bahija Zaki³, Ala-Eddine Deghmane¹, Naima Elmdaghri² and Muhamed-Kheir Taha¹*

¹ Institut Pasteur, Invasive bacterial Infections Unit, Paris, France
² Institut Pasteur du Maroc, Casablanca, Morroco
³ Laboratoire de Microbiologie, CHU Ibn Rochd, Faculté de Médecine et de Pharmacie, Casablanca, Morroco

Running Title: Meningococcal disease in Morocco

*For correspondence

Muhamed-Kheri Taha

Email mktaha@pasteur.fr

+33 1 45 68 84 38

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Meningococcal epidemiology may change unpredictably and typing of isolates of *Neisseria meningitidis* is crucial for the surveillance of invasive meningococcal disease (IMD). Few data are available of the meningococcal epidemiology in countries of North Africa. We aimed to explore the invasive meningococcal isolates from the Casablanca region in Morocco. We used whole genome sequencing (WGS) to characterize 105 isolates from this region during the period 2011–2016. Our data showed that the majority (n=100) of the isolates belonged to serogroup B. The genotyping of most of these isolates (n=62) belong to the sequence type (ST-33) of the clonal complex cc32. They also showed the same PorA and FetA markers and clustered together on the basis of WGS phylogenic analysis and seemed to correspond to an expansion of local isolates in the Casablanca region as also reported with similar isolates in several other countries in the world. These data suggest that serogroup B isolates may predominate in Morocco. They also have important impact on designating vaccination strategies.
Introduction

*Neisseria meningitidis* (the meningococcus), a Gram negative capsulated bacterium, is the causative agent of invasive meningococcal disease (IMD). IMD is characterized mainly by meningitis and/or septicemia but other clinical forms are also described (1).

*N. meningitidis* causes globally an estimated 500,000 cases of invasive meningococcal disease (IMD) per year with 10% fatality rate as well as permanent sequelae among survivors (2). Meningococci are classified into 12 serogroups based on biochemical properties of their capsular polysaccharides but 6 of them (A, B, C, W, Y and X) are responsible for almost all the disease worldwide (3). Capsular polysaccharide-based conjugate vaccines are available against serogroups A, C, W and Y while proteins-based vaccines are available against isolates of serogroup B (3).

The incidence of IMD varies worldwide with low incidence in industrialized countries between 0.3–4 cases per 100,000 population but the incidence can approach 1000 per 100,000 during epidemics in countries of the sub-Saharan Africa (4). Indeed, IMD occurs either as sporadic cases with occasional outbreaks in the industrialized countries such as in Europe. However, major epidemics occur periodically in the African meningitis belt, a region that spans countries between Senegal in the west and Ethiopia in the east (5). The distribution of serogroups also varies in space and time.

Serogroup B predominates in Europe, North America and Australasia (6). Periodic epidemics of meningococcal disease occurred in the meningitis belt that have been caused by serogroup A but serogroup A decreased in the meningitis belt since the introduction of the conjugate vaccine against isolates of serogroup A but isolates belonging to serogroups C, W and X emerged (4, 7-9). Moreover, serogroup W seems to undergo a worldwide increase since 2013 (10). Few data are available form North Africa countries (6, 11) and IMD is mainly considered as travel disease linked to the
Hajj and Umrah pilgrimages (12). Previous data from the Casablanca region in Morocco obtained between January 1992 and September 2000 showed predominance (75.5%) of serogroup B isolates with strains of the phenotype (serogroup:serotype:serosubtype) B:4:P1.15 representing 74.8% of all serogroup B isolates (13). We aimed in this work to follow the recent evolution of IMD in Casablanca region that represents 20% of the population of Morocco using whole genome sequencing (WGS) approach in order to provide more genetic resolution on the isolates and their relationship to other isolates that are currently circulating worldwide.
Materials and Methods

Ethical Statement

Isolates were obtained as part of routine activity of the Bacteriological laboratory at Hospital Ibn Rochd in Casablanca, and were analyzed anonymously for the epidemiological surveillance and control of communicable disease in the Moroccan Community. Ethical approval and informed consent were thus not required. A case of IMD was defined as the recovery of an isolate of \textit{N. meningitidis} or meningococcal DNA from a normally sterile body site, such as blood, cerebrospinal fluid (CSF), joint aspirates.

Characterization of the isolates

PCR was used to confirm suspected cases with negative culture (N=45) as previously described (14). The 105 cultured invasive isolates available from cases of IMD at the Ibn Rochd hospital for the period 2011-2016. All isolates were cultured on GCB medium plated supplemented with Kellogg supplements (15). Isolates were identified on the basis of typical morphology of colonies, Gram stain, oxidase test and API NH test (bioMerieux, Marcy l’Etoile, France), according to the manufacturer’s instructions. Antibiotic susceptibility testing was performed using Etest Minimal inhibitory concentrations (MICs) for Penicillin, amoxicillin, ciprofloxacin and Ceftriaxone, were determined by the E-test (Oxoid) methods according to manufacturers’ instructions [15] and interpreted according to the latest EUCAST guidelines, as previously recommended and described for \textit{N. meningitidis} (16).
The isolates were sent to the Institut Pasteur, Paris, France for WGS. Serogrouping and genogrouping were performed as previously described (14). Genomic DNA was extracted using the Roche MagNA Pure 96 system (Roche, Pleasanton CA 9, USA). Sequencing was performed by Illumina HiSeq 2000 sequencer (Illumina, San Diego, CA, USA) and assembled as previously described (17). Sequences are available through the PubMLST database which runs on the Bacterial Isolate Genome Sequence Database (BIGSdb) platform (18). WGS data were analyzed on the PubMLST to extract sequence type (ST) and clonal complexes according to the multilocus sequence typing analysis. Other alleles corresponding to genes of interest for antibiotic susceptibility were also extracted. WGS data were also analyzed using a “gene-by-gene” approach available through the PubMLST Genome Comparator tool using *N. meningitidis* core genome MLST (cgMLST) v1.0 that includes 1605 core loci (18). SplitsTree4 (version 4.13.1) was used to visualize the resulting distance matrices as Neighbor-net networks (19). The IDs of all these isolates are given in the supplementary Table to allow retrieving of WGS sequence in FASTA formats.
Results

Description of cases of IMD

In Morocco, notification of clinically suspected cases of IMD should be reported to the regional health authority and Case notification forms are sent to the Department of Epidemiology, Ministry of Health. The annual incidence is 2 to 3.6 per 100,000 inhabitants with 700-850 suspected cases per year fro the entire country. The Ibn Rochd University Hospital usually receives cases from the Casablanca region (about 20% of the entire country population). The surveillance for IMD is performed under the national recommendations mentioned above. Cultured bacterial isolates and samples can be sent to the Institut Pasteur of Morocco in Casablanca for identification and typing.

One hundred and forty five suspected meningococcal cases were reported and confirmed by cultured and/or PCR in Casablanca between 2011 and 2016. These cases were estimated to correspond to 25% to 40% des suspected reported cases in Casablanca. Of these 145 biologically confirmed cases, 40 (27.6%) were primary clinical materials (CSF) with no positive culture and were confirmed by PCR and 105 (72.4%) of viable cultured isolates. Most of the isolates (71%) were from CSF and 29% from blood.

We analyzed 105 isolates from Casablanca between 2011 and 2016. The age of patients ranged from 3 day to 61 years the median age is 3 years. The majority of isolates belonged to serogroup B (100 isolates; 95% for the entire period). One isolate was of serogroup C and two isolates of serogroup W and two isolates of serogroup Y. The annual distribution of isolates during this period according to serogroups is depicted in Fig.1. We detected 19% of the isolates with reduced susceptibility to penicillin G (0.12 mg/L ≤ MIC ≤ 0.25 mg/L).
Analysis of the genetic markers of the isolates

Data on the MLST were extracted from WGS and isolates were grouped into clonal complexes (Fig. 2A). Twenty-five different sequence types (STs) were identified among the isolates that were grouped into 10 clonal complexes (cc). However, 78 isolates (74% of all tested isolates) belonged to one clonal complex, the cc32. ST-33, that belonged to the cc32, was the most frequent ST (n=62 isolates) in the cc32 in the collection tested in Casablanca. Few isolates differed from ST-33 by one locus (single locus variants) or by two loci (double locus variants) (Fig. 2B).

The isolates harbored several alleles of penA encoding the penicillin binding protein2 (PBP2). Alterations of PBP2 were directly linked to the reduction of susceptibility to penicillin G in meningococci (20). Four highly related alleles (penA1, penA2, penA3 and penA22) were found in 85 isolates that were previously described to correspond to the penicillin G-susceptible isolates (20). Twenty isolates harbored diverse and altered alleles (penA9, penA19, penA33, penA35, penA217, penA295, penA709 and penA710) were previously described to correspond to isolates with reduced susceptibility to penicillin G. Indeed, the isolates harboring modified penA alleles correlated with MIC values of penicillin G.

There were several fines types (the combination of variable regions VR1 and VR2 of PorA and the variable region of FetA) but the finetype P1.19,15:F5-1 was the most frequent and was represented by 58 isolates that were all of serogroup B and cc32. These isolates were present throughout the study period (2011-2016).

The unique serogroup C isolate belonged to the ST-3327 of the cc865. The two serogroup Y isolates were different and belonged to ST-1627 (cc167) and ST-1466.
The two serogroup W isolates were of ST-11 and ST-13257 that both
belonged to the cc11 as ST-13257 was single locus variant of the ST-11.

Genomic analysis of isolates

To describe the relationships of the isolates, all the 105 cultured isolates (for the
period 2010-2016) were subjected to cgMLST analysis using the BIGSdb tools in the
PubMLST database (see Materials and Methods). The isolates grouped into several
lineages. The major cluster corresponded to the cc32 isolates that were quite separated
from the other serogroup B isolates (Fig. 3A). The isolates of serogroup
B:P1.19,15:F5-1:cc32 were highly related, we therefore compared by cgMLST the
relationships of these isolates from Morocco to all other isolates sharing the same
combination and that were available on the PubMLST (accessed on 26/12/2017). In
addition to the 58 isolates from Morocco, there were 69 other isolates between 1976
and 2017 and were from the UK, France, Brazil, Cuba, Italy, Slovenia, South Africa,
Canada, Denmark, Greece, Ivory Coast and Switzerland. The phylogenetic trees
showed clear clustering of the Moroccan isolates distinctly from the isolates from
other countries. Only few isolates from France and the UK were clustered with the
Moroccan isolates (Fig. 3B). Isolates from Casablanca region harbored the genes
encoding the antigens of the vaccines against serogroup B (Supplementary Table).

The serogroup W/cc11 isolates were compared to other W/cc11 isolates that we have
recently published in France and that belonged to the two major sublineages: the
“Anglo-French-Hajj” sub-lineage and “South American-UK” sub-lineage with its two
variants the “original UK strain” and the “UK 2013-strain” (21, 22). This analysis
clustered the two Moroccan isolates within the “Anglo-French-Hajj” (Fig. 3B).
Discussion

This report used WGS analysis as a tool for complete characterization of invasive meningococcal isolates from the Casablanca region of Morocco. Our data showed the majority of the isolates were of serogroup B in agreement with a previous report (13). Recent data from Tunisia also suggested that serogroup B predominates in the country between 1998 and 2013 (23). Serogroup B isolates were also reported in Algeria between 1992-2001 (24). A shift from a serogroup A to serogroup B was noted in Egypt since 1991 (25). However, the data remained scarce from North Africa with little molecular typing data.

The previous work on the isolate in Casablanca region reported the phenotype P1.15 for the isolate of serogroup B (13) that is in agreement with our findings on the typing of PorA (P1.19,15). Moreover, the WGS data were also available on for one group B/cc32 isolate from Morocco from 1994 on the PubMLST database (ID 26047) and clustered together with the recent isolates (26) (Figure 2). Our Data suggest the persistence of these isolates in the region and may further suggest a recent clonal expansion of these isolates with most of the isolates belonging to the major sequence type (ST-33) and few single locus and double locus variants. Of interest, the cgMLST analysis showed that these isolates seem to differ from related isolates reported in other countries suggesting an expansion of a local strain in the Casablanca region. These related isolates were responsible for large expansion as was reported in Cuba and Brazil and lead to vaccination campaigns (27). The availability of protein-based vaccines may provide a possible strategy to control the expansion of those isolates. These isolates harbor genes encoding the vaccine antigens. However, expression assays are required to confirm coverage (28). One limitation of our data is their
representativeness, but few exhaustive data are available due to limitations in reporting, diagnosis and typing (6).

The use of the WGS approach provide a thorough understanding of the circulating meningococcal isolates that provide powerful tools for high resolution in epidemiological surveillance and should help decision making to tailor vaccination strategies.
References


Zerouali K, Elmdaghri N, Boudouma M, Benbachir M. 2002. Serogroups, serotypes, serosubtypes and antimicrobial susceptibility of *Neisseria*


Acknowledgments

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Legends of figures

**Figure 1.** Distribution of the isolates of the study according to their serogroups and year of isolation.

**Figure 2.** (A) Distribution of the isolates of the study according to their serogroups and clonal complexes. (B) BURST analysis of the isolates of the clonal complex cc32 on the basis of the seven loci of the MLST. The most frequent sequence type (ST-33) is in the central circle (black). The middle circle (red) represents the single locus variants for the ST-33 and the outer circle (bleu) represents the double locus variant from the ST-33.

**Figure 3.** (A) Neighbor-net phylogenetic network of all invasive cultured isolates received of the study from the region of Casablanca, Morocco (2011-2016). The tree was drawn on the basis of cgMLST. The 10 clonal complexes and the corresponding serogroup of the isolates are indicated. (B) Neighbor-net phylogenetic network of all serogroup B isolates showing the finetype P.19,15:F5-1 in the PubMLST database: 58 isolates from Morocco in addition to 69 other isolates between 1976-2017 and were from the UK, France, Brazil, Cuba, Italy, Slovenia, South Africa, Canada, Denmark, Greece, Ivory Coast and Switzerland. The isolates from Morocco 1994 is indicated by an arrow (ID 26047) (C) Neighbor-net phylogenetic network of the two serogroup W isolates from our collection (black circle) B depicted within the tree of all invasive serogroup W isolates from France (period 2010-2016). Cases are classified in three groups using WGS data: “original UK strain”, “UK 2013-strain” and the Anglo-French-Hajj.
A

cc461/B
cc41/44/B
cc167/Y
cc174/Y
cc11/W
cc103/B
cc865/C
cc60/B
cc213/B

B

- Morocco
- UK
- France
- Others

C

The South American-UK strain
The UK 2013-strain
The Original UK strain
The Anglo-French-Hajj strain