

Predicted strain coverage of a new protein-based meningococcal vaccine in the Czech Republic

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ABSTRACT

Background: Recent data indicate that *Neisseria meningitidis* B strains cause about 70% of invasive meningococcal disease (IMD) cases in Europe and the availability of a vaccine effective against *N. meningitidis* B is desirable. A new protein-based MenB vaccine was licensed for use in Europe in January 2013. Meningococcal antigen typing system (MATS) was developed to predict strain coverage of this vaccine. Reports have recently been published for a European consortium, including aggregated data for the Czech Republic. The aim of this paper is to provide a detailed breakdown of MATS results for the Czech *N. meningitidis* B isolates.

Materials and methods: One hundred and eight *N. meningitidis* B isolates from IMD collected in the Czech Republic during 2007–2010 were selected. MATS analysis was done according to the method previously published.

Results: Based on MATS analysis, the overall estimate of strain coverage of the new MenB vaccine for a panel of 108 Czech *N. meningitidis* B strains is 74% (95% CI: 59–87%). Thirty-nine strains (36%) are predicted to be covered by a single antigen and 41 strains (38%) by more than one antigen. For 28 strains (26%), no antigen coverage was found.

Conclusions: MATS analysis showed that the new protein-based MenB vaccine could protect against a substantial proportion of IMD caused by *N. meningitidis* B in the Czech Republic. Continued detailed surveillance of IMD will be essential if the MenB vaccine is introduced to the country.

KEYWORDS

Neisseria meningitidis B – MenB vaccine – vaccine coverage – MATS – Meningococcal Antigen Typing System

SOUHRN

Křížová P., Musílek M., Vacková Z., Kozáková J., Claus H., Vogel U., Medini D.: Očekávané pokrytí kmenů novou proteinovou meningokokovou vakcínou v České republice

Cíl práce: Aktuální data ukazují, že *Neisseria meningitidis* B působí kolem 70% invazivního meningokokového onemocnění (IMO) v Evropě a dostupnost vakcíny účinné proti *N. meningitidis* B je žádoucí. Nová MenB vakcína byla registrovaná v Evropě v lednu 2013. Byl vyvinut nový systém typizace meningokokových antigenů (MATS) s cílem určit pokrytí kmenů touto vakcínou. Recentně byla publikována data pro evropské konsorcium včetně agregovaných dat pro Českou republiku. Cílem tohoto sdělení je prezentace detailních výsledků MATS pro české izoláty *N. meningitidis* B.

Materiál a metodiky: Bylo vybráno 108 izolátů *N. meningitidis* B získaných z IMO v České republice v období 2007–2010. Metoda MATS byla provedena podle publikovaných metod.

Výsledky: Podle analýzy MATS pokrytí všech českých 108 izolátů *N. meningitidis* B je 74% (95% CI: 59–87%). Bylo pozorováno, že 39 kmenů (36%) je pokryto jedním antigenem, zatímco 41 kmenů (38%) je pokryto více než jedním antigenem. U 28 kmenů (26%) nebylo zjištěno pokrytí žádným antigenem.

Závěry: Analýza MATS ukázala, že nová MenB proteinová vakcína může v České republice chránit proti podstatné části IMO způsobených *N. meningitidis* B. Pokud v České republice bude MenB vakcína zavedena, je nezbytné pokračování detailní surveillance IMO.

KLÍČOVÁ SLOVA

Neisseria meningitidis B – MenB vakcína – pokrytí vakcínou – MATS – Meningococcal Antigen Typing System

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INTRODUCTION

Invasive meningococcal disease (IMD) is one of the most devastating infectious diseases causing a high case fatality rate at times of advanced medical care. Thanks to the development and implementation of conjugate vaccines against meningococcal serogroups A, C, Y, and W135, the incidence of IMD cases caused by serogroup C has decreased in Europe and recently, serogroup B has become the leading cause of IMD. *Neisseria meningitidis* B strains have been repor-

ted to cause about 70% of IMD cases in Europe [1] and about 40–50% of IMD cases in the USA [2–4]. For this reason, the availability of a vaccine effective against *N. meningitidis* B (MenB vaccine) is desirable. However, the most significant challenges are the diversity of *N. meningitidis* B population and poor immunogenicity of serogroup B capsule. Various attempts have been made to develop a MenB vaccine effective against the whole *N. meningitidis* B population and reverse vaccinology represents a new approach [5–7]. This method

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started by whole genome sequencing of *N. meningitidis* and more than 600 antigen candidates were identified. The final version of the new four-component MenB vaccine (4CMenB) combines three primary recombinant components: fHbp (factor H binding protein) fusion protein, NadA (Neisserial adhesin A) and NHBA (Neisserial Heparin Binding Antigen) fusion protein, and the outer membrane protein PorA P1.4 of the strain causing the New Zealand outbreak. This new MenB vaccine (BEXSERO) was licensed for use in Europe in January 2013.

Meningococcal antigen typing system (MATS) ELISA was developed to predict meningococcal strain coverage of the protein-based meningococcal vaccine 4CMenB - Figure 1 [8]. MATS is considered as correlated with bactericidal activity and allows typing of large panels of strains.

MATS was used to test a collection of *N. meningitidis* B isolates, mostly from the epidemiological year 2007/2008, from seven European countries: Germany, France, England & Wales, Italy, Norway, Spain, and the Czech Republic. The predicted coverage was 78% (95% CI 63–90), ranging from 73 to 87% in individual countries [9]. The aim of this paper is to provide a detailed breakdown of the MATS results for the Czech *N. meningitidis* B isolates.

MATERIALS AND METHODS

Neisseria meningitidis strain selection

One hundred and eight *N. meningitidis* B strains collected in the Czech Republic during 2007–2010 were selected. The original collection included 120 *N. meningitidis* B isolates but 12 strains that failed to grow in the laboratory were excluded from this study. The original collection comprised all serogroup B isolates from IMD cases obtained by the National Reference Laboratory for Meningococcal Infections of the National Institute of Public Health, Prague, Czech Republic within the above mentioned period, i.e. 69 CSF isolates, 46 blood culture isolates, four isolates from other sites, and one isolate from a nasopharyngeal swab.

Neisseria meningitidis strain characterisation

Serogrouping was performed by slide agglutination, using commercial antisera. PorA variable region typing was done by sequencing, according to the Neisseria Sequence Typing Home Page [10]. Multilocus sequence typing (MLST) was done in accordance with the pubMLST neisseria sequence typing database guidelines [10]. Serogroup, PorA variable regions, and MLST were determined by the National

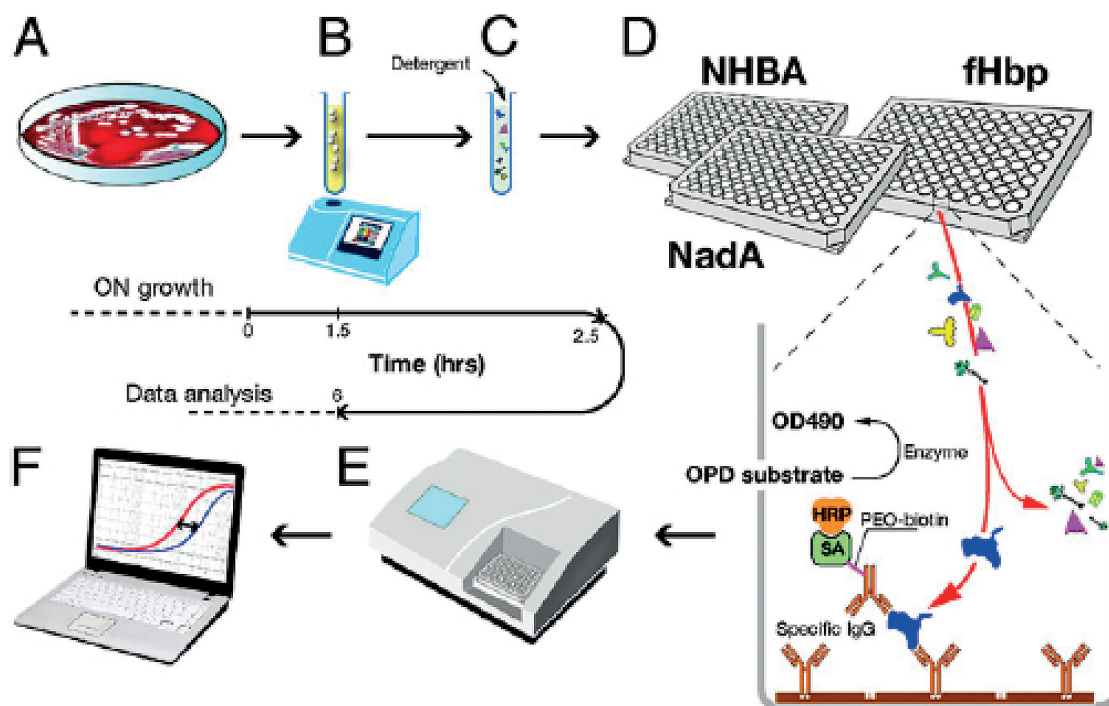


Fig 1. Schematic of the MATS ELISA method

(A) MenB bacteria are grown overnight on chocolate agar. (B) A suspension of bacteria taken from the plate is prepared to a specific OD600. (C) detergent is added to the suspension to extract the capsule and expose the antigens. (D) Serial dilutions of extract are tested in the MATS ELISA. A specific capture antibody (yellow) binds one of the antigens (example: fHbp, blue) from the extract, which is then detected with a specific biotin-labeled antibody (yellow and purple) and a streptavidin-enzyme conjugate (green and gold). (E) Plates are read at 490 nm in an ELISA reader. (F) Results are calculated by comparing the curve of OD490 vs. dilution obtained with the serially diluted unknown strain to a serially diluted reference strain tested in the same ELISA plate.

Obr. 1. Schematické znázornění metody MATS ELISA [8]

(A) Bakterie meningokoka B se kultivují přes noc na čokoládovém agaru. (B) Přípraví se bakteriální suspenze odpovídající zadané optické densitě OD600. (C) Do suspenze se přidá detergent, jehož působením se extrahuje pouzdro a dojde k uvolnění antigenů. (D) Sériová ředění extraktu se testují pomocí typizačního systému MATS ELISA. Specifická vycitávací protilátka (žlutá) na sebe naváže jeden z antigenů (například fHbp, modrá) z extraktu, který je následně detekován specifickou protilátkou značenou biotinem (žlutá a fialová), na kterou se naváže konjugát streptavidinu s enzymem (zelená a zlatá). (E) Destičky se odečtou při 490 nm pomocí čtečky ELISA. (F) Provede se výpočet výsledků na základě porovnání křivky pro OD490 proti ředění sériově naředěného neznámého kmene se sériově naředěným referenčním kmenem testovaným na stejné destičce.

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Reference Laboratory for Meningococcal Infections of the Czech Republic.

Meningococcal antigen typing system (MATS) ELISA
 MATS analysis was done according to the method published previously [8]. MATS relative potencies for fHbp, NHBA, and nadA were calculated by comparison with the *N. meningitidis* B reference strains. The predicted strain coverage was defined as the proportion of strains with MATS relative potencies greater than the positive bactericidal thresholds (PBTs) [8, 9]. Novartis supplied with the MATS methodology the University of Wurzburg, Germany, which performed MATS testing of the Czech *N. meningitidis* B isolates. Quality-assured data from the laboratory of the University of Wurzburg were collected by Novartis, subjected to further quality control, and strains with complete (4 antigens) and valid data (according to MATS acceptance criteria, consistency of documentation and QA of the originating laboratory) were selected for analysis.

Statistical analysis

95% Confidence Intervals for the positive bactericidal thresholds were estimated on the basis of MATS precision (reproducibility) determined during inter-laboratory standardization [11]. CIs do not take into account the number of strains typed.

RESULTS

Based on MATS analysis, the overall estimate of strain coverage for all 108 *N. meningitidis* B Czech strains is 74% (95% CI: 59–87%). When considering strains predicted to be covered (i.e. those that express levels of any 4CMenB antigen above the PBT or harbour PorA VR2 4), a single antigen coverage is predicted for 39 strains (36%) and 41 strains (38% of the total Czech collection spanning 2007–2010) are predicted to be covered by more than one antigen – Figure 2. Thus, the redundancy ensured by the strategy of targeting bacteria with antibodies to multiple antigens may provide an additional measure of protection against antigen mutation or loss. The distribution of PorA VR2 subtypes showed that P1.15 and P1.14 VR2 subtypes predominated in 2007–2010 (both 19%). A single strain expressed PorA P1.4 subtype, the immunodominant protein antigen contained in the OMV, and is considered covered by 4CMenB for the purpose of coverage estimation.

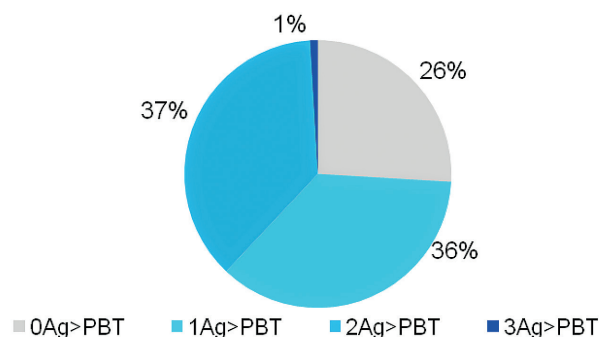


Fig 2. Coverage of *N. meningitidis* B strains by number of 4CMenB vaccine antigens, Czech Republic, 2007–2010
Obr. 2. Pokrytí kmenů *N. meningitidis* B vakcínou 4CMenB podle počtu antigenů, Česká republika, 2007–2010

Table 1. Coverage of *N. meningitidis* B strains by antigen combination^a, Czech Republic, 2007–2010

Tabulka 1. Pokrytí kmenů *N. meningitidis* B vakcínou 4CMenB podle kombinací antigenů^a, Česká republika, 2007–2010

Antigen combination ^b	Number of Strains	Percentage
fHbp+NHBA+PorA	1	1%
fHbp+NHBA	36	33%
fHbp+NadA	3	3%
NadA+NHBA	1	1%
fHbp only	32	30%
NHBA only	7	6%
No Antigen	28	26%

a: Only the actual combinations observed are presented.
 b: The antigen combinations listed are expressed at levels above the associated PBT.
 a: Uvedeny jsou pouze zjištěné kombinace.
 b: Uvedené kombinace antigenů jsou exprimovány v hladině přesahující spojený pozitivní baktericidní práh (PBT).

The coverage of strains in the panel based on the expression of fHbp, NadA, or NHBA above the MATS PBT or PorA P1.4 was then analyzed. Next, the nature of the combinations of multiple antigens present in strains that may act as targets for bactericidal antibodies elicited by 4CMenB vaccination was analysed in greater detail. The proportions of the total (n = 108) predicted to be covered based on the presence of the listed antigen(s) are shown in Table 1. Genotyping of strains was also conducted using MLST – Table 2, a standard typing system that relies on the direct sequencing of seven housekeeping genes that encode intracellular, cytoplasmic enzymes. Although MLST has limited utility in evaluating 4CMenB coverage, it provides informa-

Table 2. Distribution of clonal complexes and coverage within each complex in the *N. meningitidis* B strains, Czech Republic, 2007–2010

Tabulka 2. Distribuce klonálních komplexů a jejich pokrytí vakcínou u kmenů *N. meningitidis* B, Česká republika, 2007–2010

Clonal complex	No. of strains (percentage of a total of 108)	Coverage predicted by MATS within each cc (95% CI)
cc-41/44	20 (19%)	90% (85–95%)
cc-32	18 (17%)	100% (100–100%)
cc18	13 (12%)	85% (77–85%)
cc-269	9 (8%)	89% (22–100%)
cc-35	8 (7%)	13% (0–63%)
cc-60	5 (5%)	20% (0–100%)
Other ^a	14 (13%)	50% (14–64%)
Not Assigned ^b	21 (19%)	76% (71–86%)

a: "Other": refers to established ccs that have less than 5 strains within each cc. These include: cc-103 (3 strains), cc-1157 (2 strains), cc-174 (2 strains), cc-213 (2 strains), cc-865 (2 strains), cc-11, cc-162, and cc-461 (one strain each).
 b: "Not Assigned": Sequence Type (ST) of strains in this collection has been identified or established, but does not belong to a clonal complex (cc)
 a: "jiný": odkazuje na ustavené klonální komplexy o méně než 5 kmenech. Jedná se o cc-103 (3 kmene), cc-1157 (2 kmene), cc-174 (2 kmene), cc-213 (2 kmene), cc-865 (2 kmene), cc-11, cc-162 a cc-461 (po jednom kmeni).
 b: "nezařazený": Sekvenční typ (ST) kmenů v tomto souboru byl určen, ale nepatří k žádnému klonálnímu komplexu (cc)

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tion that can increase our understanding of the evolution of the bacterium and of the impact of mass vaccination on carriage and prevalence of the hypervirulent lineages. The most represented clonal complex (cc) in the Czech 2007–2010 collection is cc-41/44 in which 90% of strains are predicted by MATS to be covered. An equal proportion (19%) of strains have yet to be assigned to a clonal complex; of these, 76% are predicted by MATS to be covered.

DISCUSSION AND CONCLUSION

As MATS does not account for the activity of bactericidal antibodies generated from non-PorA components of OMV or synergistic effects of multiple components of 4CMenB, this is considered to be a conservative estimate of vaccine strain coverage.

A recent investigation revealed that the MATS could underestimate 4CMenB vaccine strain coverage [12]. The authors have concluded that MATS is a conservative predictive tool. However, the EMGM (European Monitoring Group on Meningococci) recommends its use for the surveillance after implementation of the 4CMenB vaccine [13]. Considering that the performance of MATS has been established in few reference laboratories in Europe, these laboratories should provide MATS testing of selected isolates on request.

Based on MATS analysis of 108 invasive meningococcal serogroup B strains collected during 2007–2010 in the Czech Republic, an estimated 74% (95% CI: 59–87%) of strains are expected to be covered by bactericidal antibody responses to the 4CMenB vaccine. Of the strains predicted to be covered (n = 80), roughly half (41 strains) are likely to be covered by more than one antigen. Thirty-two strains (30%) were covered by the fHbp antigen only and 36 strains (33%) by combination of the fHbp + NHBA antigens. In 28 strains (26%) no antigen coverage was found.

Coverage by more than one antigen provides a measure of redundancy that ensures 4CMenB can still be effective even if one target antigen is mutated or down-regulated. MATS analysis has shown that the new MenB protein-based vaccine could protect against a substantial proportion of IMD caused by *N. meningitidis* B in the Czech Republic. Continued detailed surveillance of IMD will be essential if the MenB vaccine is introduced to the country.

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