

**COMPARATIVE GENOMIC ANALYSIS OF
VIBRIO CHOLERAE AND ITS
COLONIZATION FACTORS**

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Abstract

Vibrio cholerae, the causative agent of cholera, is endemic in Malaysia and is spread through the ingestion of contaminated food or water. The aims of this study were to develop a multiple PCR assay for differentiation of *V. cholerae* from other *Vibrio* species; to compare the efficiency of PCR assay with conventional biochemical tests and API 20E.; to characterize the strains based on their biotypes, serogroups, and virulotype by using a multiplex PCR assay; to determine the genetic relatedness of strains by using genotyping methods such as RAPD-PCR, ERIC-PCR, REP-PCR, MLVA, PFGE, MLST and MVLST; to study the virulence factors which cause colonization in different serogroups of *V. cholerae*; and the influence of host environment for colonization. Four pairs of primers were designed for differential detection of sister groups of *Vibrio* species. Strains tested were differentiated into *V. cholerae* (493/338 bp), *V. parahaemolyticus* (493/409 bp), *V. vulnificus* (493/656 bp), *Vibrio* species (493 bp), and non-*Vibrio* (no amplification) based on *pntA* and *gyrB* genes. This multiplex PCR assay was more sensitive and specific than API 20E identification assay. Another multiplex PCR assay based on *ompW*, *hlyA*, *orf* complex, *toxR*, *ctxA*, *tcpI* for *V. cholerae* biotyping, serogrouping and virulotyping was developed and tested on 43 *V. cholerae* strains. A total of 22 El Tor O1 and one O139 *V. cholerae* that harboured all virulence genes were identified. One El Tor O1 *V. cholerae* presented identical virulotype to 17 other non-O1/non-O139 *V. cholerae*, while the *tcpI* gene was detected in two non-O1/non-O139 *V. cholerae*. The 43 strains were also subtyped into 38, 40, 35, 30, 35, 38, 29 and 27 profiles by RAPD-PCR, ERIC-PCR, REP-PCR, VCR-PCR, PFGE, MLVA, MLST and MVLST, respectively with discriminatory power ranging from 0.910 to 0.996. Overall, genetic diversity of non-O1/non-O139 *V. cholerae* strains was high while some of the O1 strains were indistinguishable. However,

the unrelated strains which shared the same profiles were distinguished based on the combined analyses of the eight genotyping methods. However, each method possesses its own limitations. MLST and MVLST gave precise description of point mutation but were expensive. Overall, MLVA developed in this study remains the most suitable genotyping methods based on discriminatory ability, ease of operation, cost, timeline and data management. However, a combination of several genotyping methods may overcome the inefficiency of each single method and therefore able to distinguish unrelated strains. Finally, clinical and environmental O1 strains could colonize the mouse intestines but prolonged colonization was only observed with environmental strains which showed upregulated expression of *rtxA* and *hlyA* genes. The *tcpI*⁺ non-O1/non-O139 *V. cholerae* strain was a more efficient colonizer while the non-toxicogenic O1 *V. cholerae* could colonize the mouse intestine once the virulence genes were favourably enriched and ‘turn-on’ in the host environment. In conclusion, this study provided alternative approaches for rapid differentiation of *V. cholerae* from other pathogenic *Vibrio* species, as well as to biotype, serogroup and virulotype the *V. cholerae* strains. Regardless of serogroups, year, source and location of isolation, all the unrelated strains were distinguishable and therefore suggests a high diversity of *V. cholerae* population in Malaysia. Different traits of strains possess different colonization ability and *tcpI* gene might be the key regulator for colonization in non-O1/non-O139 *V. cholerae*. However, colonization in non-toxicogenic O1 *V. cholerae* might be facilitated once the virulence genes were ‘enriched’ in the host environment.

Abstrak

Vibrio cholerae merupakan agen penyebab kolera, adalah endemik di Malaysia dan tersebar melalui pengambilan makanan atau air yang tercemar. Terdapat beberapa tujuan dalam menjalankan kajian ini, terutamanya menghasilkan beberapa kaedah PCR untuk membezakan *V. cholerae* daripada spesies *Vibrio* yang lain serta membandingkan kecekapan PCR dengan ujian biokimia konvensional dan API 20E (suatu kit untuk pengenalan bakteria). Kajian ini juga bertujuan untuk menghasilkan suatu PCR multipleks yang boleh membezakan strain-strain dari berbagai ciri biokimia, jenis kumpulan serologi, dan gen kevirulenan. Seterusnya, perkaitan genetik antara strain disiasat dengan menggunakan kaedah genotip seperti RAPD-PCR, ERIC-PCR, REP-PCR, MLVA, PFGE, MLST dan MVLST. Selain itu, faktor kevirulenan yang menyebabkan gastroenteritidis bagi strain-strain *V. cholerae* yang berbeza kumpulan serologi juga dikaji. Keseluruhannya, empat pasang primer telah direka untuk mengesan perbezaan spesies *Vibrio* yang berlainan. PCR multipleks ini mampu membezakan *V. cholerae* (493/338 bp), *V. parahaemolyticus* (493/409 bp), *V. vulnificus* (493/656 bp), *Vibrio* species (493 bp), dan juga organisma bukan-*Vibrio* (tiada amplifikasi) berdasarkan gen *pntA* dan *gyrB*. PCR multipleks ini menunjukkan kepekaan dan kekhususan sebanyak 1.0 jika dibandingkan dengan ujian biokimia konvensional dan API 20E dalam pengesanan *V. cholerae*. Ujian biokimia konvensional juga didapati lebih berkesan dan khusus berbanding dengan API 20E. Di samping itu, PCR multipleks berdasarkan gen-gen *ompW*, *hlyA*, kompleks *orf*, *toxR*, *ctxA*, dan *tcpI* untuk membezakan *V. cholerae* telah dihasilkan. Sebanyak 22 El Tor O1 dan satu *V. cholerae* O139 yang mengandungi semua gen virulens telah dikenalpasti. Satu El Tor *V. cholerae* O1 telah menunjukkan jenis kevirulenan yang sama dengan 17 strain non-O1/non-O139. Gen *tcpI* dikesan dalam dua *V. cholerae* non-O1/non-O139. Empat puluh

tiga strain juga dibahagikan kepada 38, 40, 35, 30, 35, 38, 29 dan 27 corak dengan menggunakan RAPD-PCR, ERIC-PCR, REP-PCR, VCR-PCR, PFGE, MLVA, MLST dan MVLST, masing-masing. Kuasa pembezaan bernilai antara 0.910-0.996. Secara keseluruhan, kepelbagaian genetik strain *V. cholerae* non-O1/non-O139 adalah tinggi sementara terdapat beberapa strain O1 yang tidak dapat dibezakan. Strain-strain tertentu yang berkongsi jenis corak tersebut kemudiannya dibezakan berdasarkan analisis gabungan dari lapan kaedah genotip. Selain itu, setiap kaedah mempunyai kelemahannya. MLST dan MVLST memberikan gambaran tepat dari segi mutasi titik tetapi berkongsi tinggi. Secara keseluruhan, MLVA yang dihasilkan dalam kajian ini masih merupakan kaedah yang paling sesuai untuk menaipkan genotip *V. cholerae* berdasarkan kemampuan diskriminasi, kemudahan operasi, kos, masa dan pengurusan data. Walaubagaimanapun, penggabungan beberapa kaedah menaip genotipik dapat mengatasi ketidakcekapan daripada sesuatu kaedah dan strain yang tidak berkaitan dari segi genetik dapat dibezakan. Sementara itu, strain O1 dari klinikal dan persekitaran menunjukkan penjangkapan usus tikus tapi penjangkapan berpanjangan hanya didapati pada strain persekitaran yang menunjukkan pengekspresan gen *rtxA* dan *hlyA* yang tinggi. Strain *V. cholerae* bukan serogroup O1/O139 yang mengandungi *tcpI* adalah agen gastroenteritis yang lebih kuat sedangkan strain O1 yang tidak toksik dapat menjangkit usus tikus selepas gen kevirulenan diperkayakan dan dimulakan dalam persekitaran perumah. Sebagai kesimpulan, kajian ini telah memberikan pendekatan alternatif untuk membezakan *V. cholerae* daripada *Vibrio* spesies lain yang berpatogenik selain daripada membezakan strain-strain *V. cholerae* yang berlainan ciri biokimia, kumpulan serologi dan kevirulenan dengan cekap. Semua strain yang tidak berkaitan dapat dibezakan dalam kajian ini dan menyarankan genetik kepelbagaian yang tinggi dalam populasi *V. cholerae* di Malaysia. Strain yang bersifat berlainan mempunyai kemampuan penjangkitan usus yang berbeza dan gen *tcpI* diyakinkan adalah salah satu

faktor di kalangan *V. cholerae* yang non-O1/non-O139 untuk menyebabkan gastroenteritidis. Sementara itu, penjajahan usus bagi *V. cholerae* O1 yang tidak berpatogenik dapat diamati selepas gen-gen kevirulenan 'diperkayakan' dalam persekitaran tuan rumah.

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LIST OF SYMBOLS AND ABBREVIATIONS

bp	base pair
cfu	colony forming unit
ddH ₂ O	double distilled water
DNA	deoxyribonucleic acid
dNTP	deoxynucleotide
EDTA	ethylenediamine-tetraacetic acid
ERIC	enterobacterial repetitive intergenic consensus
IMR	Institute for Medical Research
kb	kilobase
LB	Luria Bertani
M	molar
MgCl ₂	magnesium chloride
min	minute
ml	millilitre
MLVA	multiple-locus variable-number of tandem repeat analysis
mM	milimolar
NaCl	sodium chloride
ng	nanogram
OD	optical density
PCR	polymerase chain reaction
PFGE	pulse-field gel electrophoresis
RAPD	random amplified polymorphic DNA
REP	repetitive extragenic palindromic

rpm	revolutions per minute
RT	room temperature
<i>Taq</i> DNA polymerase	<i>Thermus aquaticus</i> DNA polymerase
TBE	Tris-borate EDTA
TCP	toxin-coregulated pili
TE	Tris-EDTA
Tris-HCl	Tris-hydrochloric acid
T _m	melting temperature
U	unit
UV	ultraviolet
V	volt
VCR	<i>V. cholerae</i> repeats
VPI	<i>Vibrio cholerae</i> pathogenicity island
WHO	World Health Organization
w/v	weight/volume
~	approximately
°C	degree Celcius
=	equals to
<	less than
>	more than
%	percent
μl	microlitre
μg	microgram
μM	micromolar