Acquisition of carbapenemase-producing Enterobacteriaceae by healthy travellers to India, France, February 2012 to March 2013

E Ruppé (etienne.ruppe@gmail.com) 1,2,3, L Armand-Lefèvre 1,2, C Estellat 1,2, A El-Mniai 1,5, Y Boussadia 1,5, P H Consigny 1,6, P M Girard 7, D Vittecoq 1,6, O Bouchaud 1,6, G Pialoux 9, M Esposito-Farèse 4,5, B Coignard 8, J C Lucet 2,3,12, A Andremont 1,2,3, S Matheron 3,13

1. AP-HP, Hôpital Bichat, Laboratoire de Bactériologie, Paris, France
2. INSERM, IAME, UMR 1137, F-75018 Paris, France
3. Univ Paris Diderot, IAME, UMR 1137, Sorbonne Paris Cité, Paris, France
4. AP-HP, Hôpital Bichat, Département d’Épidémiologie et Recherche Clinique, URC Paris-Nord, Paris, France
5. INSERM, CIC 1425-EC, Paris, France
6. Institut Pasteur, Centre Médical, Centre d’ Infectiologie Necker-Pasteur, Paris, France
7. AP-HP, Hôpital Saint-Antoine, Maladies Infectieuses et tropicales, Paris, France
8. AP-HP, Hôpital de Bicêtre, Maladies Infectieuses et Tropicales, Le Kremlin-Bicêtre, France
9. AP-HP, Hôpital Avicenne, Maladies Infectieuses et Tropicales, Bobigny, France
10. AP-HP, Hôpital Tenon, Maladies Infectieuses et Tropicales, Paris, France
11. Institut de Veille Sanitaire, Saint Maurice, France
12. AP-HP, Hôpital Bichat, Unité d’Hygiène et de Lutte contre les Infections Nosocomiales, Paris, France
13. AP-HP, Hôpital Bichat, Maladies Infectieuses et Tropicales, Paris, France

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Healthy travellers to countries where carbapenemases-producing Enterobacteriaceae (CPE) are endemic might be at risk for their acquisition, even without contact with the local healthcare system. Here, we report the acquisition of CPE (two OXA-181, one New Delhi metallo-beta-lactamase 1 (NDM-1)) in three healthy travellers returning from India. The duration of CPE intestinal carriage was less than one month. The results indicate that healthy travellers recently returning from India might be considered as at risk for CPE carriage.

We report the acquisition of carbapenemases-producing Enterobacteriaceae (CPE) in three healthy French travellers returning from India, who declared no contact with any local healthcare centres during their journey in this country.

Healthy travellers carrying carbapenemase-producing Enterobacteriaceae
As part of the VOYAG-R study (ClinicalTrials.gov Identifier: NCT01526187), volunteers who had planned to travel to intertropical areas for a three-day to three-month trip were recruited in six centres for travel vaccinations in the Paris area, from February 2012 to March 2013. In total, 574 travellers (222 men and 352 women) were included, who visited 72 intertropical countries located in the Americas (n=183 travellers), Africa (n=195) or in the Middle East and south-east Asia (n=196). This included 57 travellers who visited India. If travelling in groups, only one self-designated volunteer was solicited. Included travellers were those with a negative stool sample for multidrug-resistant Enterobacteriaceae (MRE) during the week preceding departure and those who provided a stool sample within a week after their return. Each sample was accompanied with a self-completed questionnaire. Before departure the traveller informed on the dates of departure and return, the visited country, the number of accompanying travellers, the malaria prophylaxis, and the type and purpose of the travel. After return, the traveller reported on the occurrence of digestive disorders during the travel, the intake of antibiotics, any contact with the healthcare system at travel destination and the compliance with malaria prophylaxis. During the follow-up the traveller informed of any antibiotic intake and purpose, any hospitalisation and any new travel to intertropical countries. If positive for MRE after return, the traveller was asked to provide stool samples one, two, three, six and 12 months after return, until no MRE could be detected. Among 57 travellers who had visited India, three returned to France with CPE intestinal carriage (Table).

Traveller 1 (C4-049)
A woman in her early fifties, had travelled alone as a backpacker and tourist to India for 17 days in April 2012. Upon return, she did not report any digestive disorders, any antibiotic intake or any contact with the local healthcare system during her travel. Investigation of stool samples revealed four phenotypically distinct Escherichia coli, including one that produced both a CTX-M group 1 and an OXA-181 carbapenemase. One
month after return, a CTX-M group 1-producing *E. coli*, which displayed a different resistance pattern to that of the *E. coli* recovered at return, was also detected. Two months after return, a stool sample from traveller 1 was negative for MRE.

**Traveller 2 (C4-417)**
A woman in her late twenties, travelled to northern India for 10 days in November 2012, with another person on a tour. She did not report any digestive disorders, any antibiotic intake or any contact with the local healthcare system during her travel. Direct cultures of stool samples collected at her return on agar media were negative, but cefotaxime enrichment broth yielded a CTX-M group 1-producing *E. coli*. Furthermore, the carbapenemase specific enrichment procedure used for this study yielded an OXA-181-producing *E. coli*. Traveller 2’s stool sample, originating from one month after return, was negative for any MRE carriage.

**Traveller 3 (C4-422)**
A woman in her early thirties, travelled on her own to southern India for one month in January 2013, where she alternatively backpacked, participated in touristic tours and visited relatives living in India. At return, she reported having experienced digestive disorders, but she had not taken any antibiotics nor visited any healthcare centre during her journey in the country. From her stool sample at return, six phenotypically distinct *E. coli* were identified, among which one produced both CTX-M group 1 and New Delhi metallo-beta-lactamase 1 (NDM-1) carbapenemase. At months 1 and 2 after return, she was no longer carrying any CPE, but was still carrying one CTX-M group 1-producing *E. coli*. A stool sample from three months after return was negative for MRE.

**Laboratory investigations**

**Detection of multidrug-resistant Enterobacteriaceae**
Stool samples were stored at room temperature by the traveller until shipped by postal services to the Bacteriology laboratory of the Bichat-Claude Bernard Hospital, Paris, France, where they were cultured immediately upon reception. Approximately 10mg of stool was plated onto a chromID extended-spectrum beta-lactamases (ESBL) agar media (bioMérieux, Marcy-l’Etoile, France) and onto a bi-valve ESBL agar (AES Chemunex, Ivry-sur-Seine, France). In parallel, approximately 100mg of stool was diluted in 10mL of brain heart infusion (BHI) broth, of which 1mL was diluted to a BHI broth supplemented with 1.5mg/L cefotaxime and another 1mL to a BHI broth supplemented with 0.5mg/L ertapenem, and incubated overnight, until 100µL of each broth were respectively plated onto a chromID ESBL agar media and a Drigalski agar plate with disks of ertapenem and imipenem, as described [1]. Plates were incubated 48h at 37°C in aerobic conditions. All colony-forming units (CFUs) with distinct morphotypes on chromID ESBL agar media and CFUs growing within the normal inhibition radius of carbapenem disks (www.sfm-microbiologie.org) were further identified by mass spectrometry (MALDI Biotyper, Bruker, Bremen, Germany) and tested for antibiotic susceptibility by the disc diffusion method, as recommended by the French Society for Microbiology (www.sfm-microbiologie.org).

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**Table**

<table>
<thead>
<tr>
<th>Traveller ID</th>
<th>Strain</th>
<th>Species</th>
<th>Beta-lactamases</th>
<th>Co-resistances</th>
<th>Return</th>
<th>Follow-up</th>
</tr>
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<tbody>
<tr>
<td>C4-049</td>
<td>C4-049Ec1</td>
<td><em>Escherichia coli</em></td>
<td>CTX-M group 1</td>
<td>TE</td>
<td>Month 1</td>
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<td>C4-049</td>
<td>C4-049Ec2</td>
<td><em>Escherichia coli</em></td>
<td>CTX-M group 1</td>
<td>FQ, SXT, TE</td>
<td></td>
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</tr>
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<td>C4-049Ec3</td>
<td><em>Escherichia coli</em></td>
<td>CTX-M group 1</td>
<td>FQ, TE</td>
<td></td>
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<td>C4-049</td>
<td>C4-049Ec4</td>
<td><em>Escherichia coli</em></td>
<td>OXA-181 and CTX-M group 1</td>
<td>FQ</td>
<td></td>
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<tr>
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<td>C4-049Ec5</td>
<td><em>Escherichia coli</em></td>
<td>CTX-M group 1</td>
<td>GM, FQ, SXT, TE</td>
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<td>FQ, TE</td>
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<td>C4-417Ec2</td>
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<td>OXA-181</td>
<td>FQ</td>
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<td><em>Escherichia coli</em></td>
<td>CTX-M group 1 and pAmpC</td>
<td>GM, FQ, TE</td>
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<tr>
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<td>FQ, TE</td>
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<td>C4-422Ec4</td>
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<td>CTX-M group 1</td>
<td>FQ, SXT, TE</td>
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<tr>
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<td><em>Escherichia coli</em></td>
<td>pAmpC</td>
<td>FQ, SXT, TE</td>
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<td><em>Escherichia coli</em></td>
<td>NDM-1 and CTX-M group 1</td>
<td>FQ, AN, GM, SXT, TE</td>
<td></td>
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</tr>
</tbody>
</table>

AN: amikacin; FQ: fluoroquinolone; GM: gentamicin; ID: identifier; NDM-1: New Delhi metallo-beta-lactamase 1; pAmpC: plasmid-encoded AmpC-type cephalosporinase; SXT: trimethoprim/sulfamethoxazole; TE: tetracycline.

Black-filled cells indicate the detection of the strain in question, grey cells indicate that the strain was not detected, light grey cells indicate that no stool sample was requested (i.e. follow-up completed).
Characterisation of the resistance mechanisms

Total DNA of MRE was extracted by the EZ1 DNA Tissue Kit processed on the EZ1 instrument (Qiagen, Courtaboeuf, France). blaCTX-M, blaTEM, blaSHV, blaOFC, blaVIM, blaIMP and blaOXA-48 were targeted with specific polymerase chain reaction (PCR) primers, as described [1-3]. blaNDM was targeted using the following primers: NDM-F 5’-CTGAGCACCCAATTACCGG-3’ and NDM-R 5’-CGTATGAGTGATGGCGCCG-3’. Plasmid-encoded AmpC-type cephalosporinases (pAmpC), blaCTX-group, blaINT-group, blaMOX and blaOX, were targeted by the wide-range AmpCU-F 5’-GCARACSCTGTTYGAGMTDGG-3’ and AmpCU-R 5’-CTCCCARCCYARYCCCTG-3’ primers. Amplicons of carbapenemases-encoding genes were Sanger-sequenced with the Big Dye terminator v3 kit (Applied Biosystems, Courtaboeuf, France) for final identification.

Ethical issues

The VOYAG-R study was approved by the ‘Comité de Protection des Personnes’ Ile de France IV (14 November 2011).

Discussion and conclusion

MRE that produce ESBL and/or plasmid-encoded AmpC-type cephalosporinases (pAmpC) have spread massively in developing countries. This phenomenon likely results from suboptimal hygiene living conditions and uncontrolled antibiotic usage [4]. Therefore, travellers may be at risk for MRE acquisition when visiting countries in which the MRE prevalence is high. In recent years, studies focusing on the acquisition of MRE during travel abroad have shown that MRE acquisition rates ranged from 14.0% to 30.5% [5-10]. Surprisingly, in those studies no CPE was isolated from healthy travellers, despite them having visited CPE endemic areas such as the Indian subcontinent. Some sporadic cases of CPE importation, with no connection with any healthcare centre, have been reported, all for travellers returning from India [11-13], but not healthy travellers.

We report the acquisition of CPE in three healthy French travellers returning from India, who declared no contact with any healthcare centres in this country. These findings are worrisome as they attest to the development of a community reservoir for CPE, at least in India.

On a positive note, the duration of CPE carriage in the three travellers was less than one month. In former studies of acquisition of MRE by travellers, it was not clear whether the MRE carriage could be short [5] or long [7,9]. Despite the limited number of acquisitions of CPE, our results might suggest that, travellers immediately returning from CPE endemic areas should be considered at risk for CPE carriage, while in the absence of antibiotic exposure, travellers at several months after their return might not pose such a risk.

Our results stress the need for a specific cultivation method for assessing the intestinal carriage of CPE when suspected, such as the one we used [1], because some CPE such as those producing OXA-48 type carbapenemases (including OXA-181) do not grow on agar media formulated to detect ESBL-producing Enterobacteriaceae [1]. Some CPE might have been missed in former studies because no specific and sensitive detection of CPE was used.

In conclusion, we report here the acquisition of CPE by healthy travellers to India without contact with any local healthcare centre, while in this country. In addition to repatriated patients or patients who have recently been hospitalised abroad, travellers may be considered at occasional risk for CPE carriage.

Acknowledgments

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Conflict of interest

None declared.

Authors’ contributions

ER, LAL, CE, BC, JLM, AA and SM designed the study, analysed the data, and reviewed the manuscript. YB, PHC, PMG, DV, OB, GP and SM included the travellers. AEM performed the microbiological analysis. MEF and CE performed the statistical analysis. ER wrote the manuscript.


