Cell Specificity of *Helicobacter pylori* Cytotoxin Is Determined by a Short Region in the Polymorphic Midregion

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There are two alleles of the vacuolating cytotoxin gene from *Helicobacter pylori*, which code for toxins with different cell specificities. By analyzing the phenotypes of natural and artificial chimeras between the two forms of the protein, we have delimited a short stretch of amino acids which determine the cell specificity.

*Helicobacter pylori* produces a potent excreted cytotoxin, VacA, which causes a massive vacuolar degeneration in the target cells in vitro and gastric epithelial erosion in vivo (4, 13). VacA is found as high-molecular-weight oligomeric structures of an 87-kDa polypeptide with either six- or sevenfold radial symmetry (9, 10). Each monomer can be cleaved proteolytically at a specific site into two fragments of approximately 37 and 58 kDa that remain associated in the holotoxin (14). Biological and structural data suggest that VacA is similar to the AB family of dichel toxins, which consist of two distinct moieties, A and B, involved in toxic activity and membrane interaction, respectively (5, 11, 14).

VacA binds to the surfaces of target cells and then is translocated to the cytosol, where it is active (6, 7). Intracellular expression of a transfected truncated *vacA* gene from which most of the sequence coding for the 58-kDa subunit has been deleted results in cell vacuolation, indicating that 37-kDa toxin subunit is responsible for the vacuolating activity (6). More recently, Reyrat et al. (14) have shown that the 58-kDa subunit is involved in binding to the target cell. Toxicity has been associated with mosaicism in *vacA* genes (1, 5). Two variants of the middle region of the gene, m1 and m2, have been described (1). Most isolates with the m1 form are toxic to HeLa cells, whereas the m2 forms are essentially nontoxic to these cells. However, both forms are toxic to primary cultured human gastric cells and the rabbit kidney epithelial cell line RK-13 (11). Hence, the m2 form of VacA is fully toxic if the appropriate cells are used, which is consistent with the lack of statistical correlation between *vacA* allele or cytotoxicity to HeLa cells and disease (8) and the high incidence of peptic ulcer and gastric cancer found in the Chinese population, where the m2 allele is prevalent (12). We have previously shown that the absence of activity of the m2 cytotoxin in HeLa cells is due to a lack of interaction with the cell, indicating that the two forms of the toxin differ in their binding domains (11).

The major difference between the two types of protein is in an approximately 300-amino-acid region in the 58-kDa subunit (1, 5), and it is hence likely that it is this difference which determines the cell specificity of the toxin (11). In order to demonstrate this and to delimit the functional differences more precisely, we have analyzed the toxic activities of the products of natural and artificial chimeric *vacA* genes in which different parts of the midregion of an m1 gene have been replaced by the corresponding regions from an m2 gene.

Recently, unusual *vacA* gene hybrids (m1-type proximal and m2-type distal) were identified (2, 12). We have sequenced the complete *vacA* gene from a naturally chimeric m1-m2 *H. pylori* strain, ch2 (GenBank accession no. AF191639) (2). The predicted product of this gene is highly similar to the m1 form of the protein up to amino acid 648 and is highly similar to the m2 form from amino acid 657 onwards (Fig. 1). Both water extracts and purified cytotoxin from ch2 induced vacuolation and neutral red uptake in HeLa cells to levels similar to those induced by a canonical m1 toxin from strain G27 (Fig. 2A). A canonical m2 toxin from strain 95-54 (11) failed to induce vacuolation in this assay but was fully toxic to RK13 cells (Fig. 2C). Hence, the chimeric ch2 toxin has the m1 toxic phenotype, indicating that the midregion from amino acid 657 onwards plays no role in the phenotypic difference between the m1 and m2 forms.

The above result indicates that the first 148 amino acids of the midregion between amino acids 501 and 657 contain the determinants of toxicity to HeLa cells. To demonstrate this, we have engineered the *vacA* gene in m1 strain G27 in order to replace this region with the corresponding region from the m2-type *vacA* gene from strain 95-54. In addition, we have dissected this region further by two chimeric *vacA* genes in which the sequences coding for amino acids 496 to 535 and 535 to 696 have been replaced by the corresponding m2 sequences.

The strategy used to generate the chimeric *vacA* genes in G27 is shown in Fig. 3A. First, a G27 recipient strain was created by natural transformation with a plasmid construct containing the *vacA* gene sequences from m1 strain CCUG17874 (14) coding for the 37-kDa subunit, followed by a kanamycin resistance cassette, followed by about 1,000 bp of sequence immediately downstream of the stop codon (GenBank accession no. AF191640). Hence, in the resulting kanamycin-resistant strain (*H. pylori* G27:Py) the *vacA* gene sequences coding for the whole of the 58-kDa subunit and outer membrane exporter have been replaced by the kanamycin resistance cassette. Chimeric midregions were created in a delivery plasmid, PQE30Delivery (GenBank accession no. AF191638), which contains the complete CCUG17874 *vacA* gene, followed immediately by a chloramphenicol resistance gene, followed by *vacA* gene 3′ flanking sequences (3). The *vacA* gene sequences in this plasmid between the unique...
EcoNI and AflII sites were replaced by PCR products corresponding to the chimeric midregion. The PCR products were prepared in two steps. First, the parts of the m1 and m2 genes were amplified using primers based on their respective sequences. For each construct, the 3' primer on the left part was complementary to the 5' primer of the right part. The products of these PCRs were purified and mixed in a second reaction containing only the leftmost and rightmost primers such that a product could be obtained only if the complementary regions of the two fragments annealed through the overlapping complementary sequences. The resulting chimeric delivery constructs were transformed into the recipient strain G27::Pγ, and gene replacement strains PJ2 (GenBank accession no. AF191637), PIV6 (GenBank accession no. AF191636), and PAB25 (GenBank accession no. AF191635) were identified by resistance to chloramphenicol and sensitivity to kanamycin. The products of the constructs are shown schematically in Fig. 3B.

Water extracts from these strains were prepared, and the presence of VacA was determined by immunoblot analysis using rabbit sera raised against a recombinant m1 form of the protein (15). The results demonstrate that the mutant strains expressed mature 87-kDa VacA protein (Fig. 4). Laser densitometry of the filter revealed that the weakest band was 50% as intense as that of the parental G27 strain. However antibodies raised against the m1 form of the protein recognize the m2 form less well (11). The water extract of strain G27::PJ2 failed to induce vacuolation of HeLa cells (Fig. 2A). However, full toxic activity, equivalent to that of the parent G27 strain, was found when RK13 cells were used in the assay (Fig. 2C). Hence, the PJ2 chimera has the m2 toxic phenotype, thus confirming that the functional difference in toxic phenotype between the m1 and m2 forms resides in the first 160 amino acids of the midregion. A water extract from G27::PIV6 (Fig. 2B) also failed to induce vacuolation in HeLa cells but was
chimeric protein, since the same water extract was as toxic as that of the parental strain when RK13 cells were used in the assay.

These data demonstrate that the amino acid differences between the m1 and m2 forms in a short segment of the toxin at the 5’ extremity of the midregion play a major role in defining cell specificity. m2 forms of the toxin have a 21-amino-acid insert in this region which is lacking in m1 forms. This insert, however, plays no role in determining cell specificity, since insertion of these sequences into the G27 vacA gene (G27::P11) had no effect on the HeLa cell toxicity of the VacA product (Fig. 2B).

The data define a short region of 148 amino acids from the beginning of the midregion to the m1-m2 junction in strain ch2 which determines the phenotypic differences in target cell specificity between the m1 and m2 forms of the protein. Within this region, the first 35 amino acids must correspond to the m1 sequence for HeLa cell cytotoxicity. Excluding the 21-amino-acid insert, there are 13 amino acid differences between the m1 and m2 sequences used. However, an analysis of a number of genes from different geographic areas (X. Ji and J. L. Telford, unpublished data) reveals that only 10 of these positions consistently differ between the m1 and m2 forms (Fig. 1). The next 113 amino acids appear to influence the potency but do not completely eliminate toxicity to HeLa cells. In this region there are 49 amino acid differences between the genes used, of which 31 are consistently different between the m1 and m2 forms. These differences, however, do not affect the innate toxicity or functionality of the toxin, since all chimeric proteins tested were fully toxic to RK13 cells.

Pagliaccia et al. (11) have demonstrated that lack of toxicity of an m2 form of the protein correlates with lack of cell surface binding in HeLa cells and have postulated different receptors for the two forms. A key result from these studies was that the m2 form was fully toxic to HeLa cells if the lack of surface interaction could be overcome by intracellular expression of the protein. From these data, however, it is not possible to distinguish between the existence of two independent receptors, one for each form of the toxin, or a common receptor for both forms on RK13 cells and a different m1-specific receptor on HeLa cells. Hence, although it is clear that the region identified above is necessary for interaction with the HeLa cell receptor, it cannot be excluded that other regions of the protein are also involved. Furthermore, the results could also be explained if the receptor is polymorphic either in protein sequence or glycosylation such that the m2 toxin fails to recognize the HeLa cell isoform. Finally, the data raise the question of how or why the protein has evolved in two forms with such large differences over a 300-amino-acid segment if only a relatively short region determines cell specificity. These questions will require identification of the receptor(s) and an elucidation of the mechanism of interaction.

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