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Short communication

Clostridium difficile (including epidemiology)

A genetic assay for gene essentiality in *Clostridium*

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Clostridium is a bacterial genus composed of around 100 species which are either of industrial or medical importance. Arguably, the most noteworthy clostridial species are those that cause human and animal diseases, and in particular *Clostridium difficile*. Responsible for *Clostridium difficile*-associated disease (CDAD), it is the leading cause of hospital-acquired and antibiotic-associated diarrhoea worldwide. In the US, *C. difficile* was responsible for almost half a million infections and 29,000 deaths in 2011, while similar rates of infection are estimated in Europe [1,2]. Treatment options remain limited [3]. Moreover, the emergence of strains resistant to currently used antibiotics [4] has led the CDC to include *C. difficile* as one of the major "Antibiotic Resistance Threats in the United States, 2013" [5]. New therapies are required.

One option is to identify those genes and their products which are essential for the bacterium's survival, and then develop interventions

ABSTRACT

Essential genes of pathogens are potential therapeutic targets, but are difficult to verify. Here, gene essentiality was determined by targeted knockout following engineered gene duplication. Null mutants of candidate essential genes of *Clostridium difficile* were viable only in the presence of a stable second copy of the gene.

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> that target that function. Direct demonstration of gene essentiality is not, however, a straightforward process. One indirect approach, is to use high-throughput sequencing methods (Transposon-Directed Insertion Site Sequencing, TraDIS) to allow simultaneous screening of saturating transposon libraries. Those genes found not to contain transposon insertions are presumed to be essential for growth under the conditions used to make the library [6]. When applied to the C. difficile strain R20291, 404 genes were suggested to be essential [7]. Another strategy, and that adopted here, is to show that a particular gene can only be inactivated if the target cell is made a merodiploid through the addition of a second functional copy of the gene in question [8]. Our approach was made possible by the properties and characteristics of two clostridial gene tools: (i) the ClosTron [9], a Group II intron retargeting mutagen which is absolutely reliant on the presence of a specific recognition sequence within the gene to be inactivated, and (ii) Allele-Coupled Exchange (ACE) technology, which allows the rapid integration of DNA of any size or complexity into the C. difficile genome [10].

> For proof of principle studies we chose *metK* (S-adenosylmethionine synthetase) and *trpS* (tryptophan tRNA synthetase) as both have been shown to be essential in *Bacillus subtilis* [11] and were included in the list of 404 essential *C. difficile* genes identified using TraDIS [7]. Retargeted ClosTrons directed against each gene were designed and constructed using standard procedures [9]. In

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parallel, both genes were resynthesized exactly as the wildtype, except over the two 45 nucleotide regions that encompassed two predicted ClosTron target sequences. Here, several synonymous codon replacements were made, in order to change the DNA sequence without affecting the amino acid sequence of the encoded MetK or TrpS protein (Fig. 1). As a consequence, these regions were no longer recognised as intron targets using the Peruka algorithm [12]. In each case, care was taken not introduce any rare-codons into the amino acid sequence so the protein's translational efficiency would be as close as possible to the native proteins.

The ACE integration vector pMTL-JH18 [10] is designed (through provision of flanking asymmetric homology arms) to create a deletion in the *pyrE* gene (encodes orotate phosphoribosyl-transferase) which confers on the host resistance to 5-fluorouracil (FOA). The two synthetic genes including their natural promoter, were cloned into pMTL-JH18 [10], transformed into *C. difficile* 630 Δerm and plated on media supplemented with thiamphenicol. Single crossover integrants, selected on the basis of their larger colony size, were plated onto minimal media containing FOA and uracil [10]. The majority of the FOA-resistant (^R) colonies that arose (e.g. 16 of 17 in the case of *metK*) were clones in which the *metK or trpS* had integrated into the genome concomitant with inactivation of the *pyrE* locus. Authenticity was confirmed by undertaking a PCR screen employing two primers flanking *pyrE* (Fig. 2). Those few

FOA^R colonies which arose that lacked an insertion most likely represented spontaneous mutants in *pyrE* or *pyrF*.

The resulting merodiploids for *metK* and *trpS* were respectively used in parallel with the wild type $630\Delta erm$ strain, as recipients in ClosTron mutagenesis experiments directed against the native *metK* and *trpS* genes. Following the transfer of the four retargeted plasmids (two targets for each gene) to both the wild type and the appropriate merodiploid strain, putative ClosTron mutants were isolated as erythromycin-resistant (Em^R) colonies. In the case of the wild type $630\Delta erm$ strain, despite screening 70 such Em^R clones for each target, no intron insertions in either metK or trpS were detected. In contrast, intron insertions were readily found when a merodiploid strain was the recipient, with the majority of the 20 Em^R colonies tested for each target being intron insertions in either *metK* or *trpS* (Table 1). These data demonstrate that the *metK* and trpS genes are essential under the growth conditions employed and cannot be inactivated unless a second, functional copy of the gene is also present in the cell.

Having demonstrated the utility of the system, *CD0274*, a previously [13] suspected essential gene identified and annotated as *gldA* (glycerol dehydrogenase) in *Clostridium beijerinckii* and *dhaT* (1,3-propandiol oxidoreductase) in *C. difficile*, was used to validate the method. *CD0274* is proposed to play a pivotal role in the detoxification of the toxic metabolite methylglyoxal (MG) [13,14].

CD02	74 CT Target - 548a						
wt Synth	L T N S E T K D K K S I K K N S M 451 CTT ACA AAT TCA GAA ACT AAA GAT AAA TCT ATA AAG AAT AGT AT 500 451 CTT ACA AAT TCA GAA ACT AAG GAT AAg AAg AAT AAA AAA AAG AAA AAA AAA AAA AAA AAA AAA AAA AAC tcT AT 500						
wt Synth	Y A K A S I I D P E L M V TV M P 502 TAT GCA AAA GCA TCA ATA ATT GAC CCA GAA CTT ATG GTT ACA ATG CCA A 502 TAC GCt Aag GCt Agt ATt ATa GAC CCA GAA CTT ATG GTa ACt ATG CCt A 500 TAC GCt Aag GCt Agt ATt ATa GAC CCA GAA CTT ATG GTa ACt ATG CCt A 500 TAC GTa ACt ATG CCt A						
wt Synth	KHIASVGFDALAHNME551AACATATAATTGCATCAGTTGATGGACTTGCCCATAATATGGAA600551AgCAcATtATaGCtagtGTaGGtTTcGAcGCTGCCCATAATATGGAA600551AgCAcATtATaGCtagtGTaGGtTTcGAcGCACTTGCCCATAATATGGAA600						
wt Synth	A Y L S N G R N P L A D V O A I Y 601 GCA TAT CTT TCT AAT GGA AGA AAC CCT CTT GCA GAT GTA CAA GCT ATT TA 601 GCA TAT TTa agt AAc GGT AGg AAt CCA tTa GCt GAC GTt CAg GCa ATa TA 650						
wt Synth	G I E L I S E N L I K V Y N D V 651 T GGG ATT GAA CTT ATA TCA GAA AAC CTA ATA AAG GTA TAT AAT GAT						
<u>metK</u>	[↑] CT Target - 623a L V D Y L R <u>P D G V K T Q V T V E</u>						
wt Synth	501 T TTA GTT GAT TAT TTA AGA CCA GAT GGA AAA ACT CAA GTT ACT GTT GAA T 501 T TTA GTT GAT TAT TTA CGC CCt GAC GGC AAA ACA CAA GTC ACA GTC GAA T 550 Y E G S K A V R V H T V L I S A O						
wt Synth	551 AT GAA GGA AGT AAA GCT GTA AGA GTA CAT ACA GTT CTT ATA TCA GCT CAA 600 551 Ac GAA GGt AGT AAg GCc GTg CGc GTg CAT ACA GTC CTT ATT TCA GCc CAA 600						
wt Synth	601 CAT TGT GAA ACA GTA TCA AAT GAT AAA ATA AGA GAA GAT TTA ATT AAT CA 601 CAT TGE GAA ACE GTg TCA AAE GAT AAg ATA eGt GAA Gac CTt ATC AAT CA 650						
wt Synth	V I K E V I P A E L D E T K 651 T GTT ATA GGA GTT ATA CCA GCA GAA TTA CTA GAA GAA AAA A 700 651 C GTG ATC AAA GAA GCA GCA GGA GTA CTA GAA AAA A 700						
trpS	CT Target - 676s CT Target - 436a, CT Target - 460a,						
wt Synth	A D I L L Y O TYD L V P V G DYD 401 CT GCT GAC ATA TTG TTG TAT CAA ACT GAC CTA GTT CCA GTT GGA GAT GAC 401 CT GCT GAT ATe TTa TAT CAg Acg GAC tTg GTe CCA GTe GGA GAE GAt 450						
wt Synth	OKOHLELARD AN RFN 451 CAA AAA CAA CAT TTA GAA TTA GCT CGT GAT TTA GCA AAT AGA TTT AAT AA 500 451 CAA AAG CAG CAT TTG GAA TTG GCT CGT GAT TTA GCA AAT AGA TTT AAT AA 500						

Fig. 1. Protection of the synthetic merodiploid copy of the candidate essential gene. The ClosTron group II intron recognises a 45mer target site on the sense (s) or anti-sense strand (a) displayed by a box encompassing the two target sites for each candidate essential gene. The native gene (wt) is shown above, and the re-synthesised gene below (Synth). The degenerate changes made to synthetic gene codons are shown in bold lowercase. Those changes made outside of the boxed target regions represent alterative ClosTron target sites not used in the study.



Fig. 2. Construction of target gene merodiploid cell lines. (**A**) The synthetic gene (*CD0274*) lacking the intron target (dashed box) to be delivered is cloned between the two homology arms of the replication defective vector, pMTL-JH18, and integrated into the genome using ACE [10]. Concomitant with integration, the *pyrE* gene is inactivated resulted in a cell that is auxotrophic for uracil and resistance (^R) to fluoroorotic acid (FOA). Insertion of the group II intron is selected on the basis of acquisition of resistance to erythromycin due to acquisition of the activated *ermB* gene (indicated as a filled arrow). Insertion of the intron into the original CD0274 gene only takes place in the merodiploid cell and not the wild type strain. The position of the two PCR primers F1 (Cdi630:pyrD-F1) and R1 (Cdi630:CD0189-R1) used to confirm insertion of the merodiploid gene at the *pyrE* locus is shown beneath the relevant region of the genome. (**B**) PCR screening of three double crossover clones of each target gene (*CD0274, metK* and *trpS*) using primers Cdi630:pyrD-F1 (F1) and Cdi630:CD0189-R1 (R1). The molecular weight marker (MW) used (lane 1) is an NEB 1 Kbp DNA Ladder. Lane 2 on all three gels is water only negative control, lane 3 is the wildtype (expected size ~2.8 kbp). *C. difficile* 630*Δerm* DNA control, lanes 4 are candidate clones of the target genes as indicated (expected size ~ 3.3 kbp).

Table 1

ClosTron Group II intron Insertion Frequencies at Target Gene.

Target gene	ClosTron target	Wild type recipient			Merodiploid recipient		
		Colonies screened	Intron insertions	Frequency (%)	Colonies screened	Intron insertions	Frequency (%)
metK	548s	70	0	0	20	20	100
	676a	70	0	0	20	20	100
trpS	548s	70	0	0	20	20	100
	623a	70	0	0	20	18	90
CD0274	436a	70	0	0	20	15	75
	460a	70	0	0	20	20	100

Accordingly, the CD0274 gene was synthesised as before and the sequences of the two highest scoring [12] ClosTron targets altered through the use of degenerate codon sequences (Fig. 1). The synthetic gene was sub-cloned into pMTL-JH18 [10] and integrated into the genome at the *pyrE* locus using ACE [10]. Following the constructing of the CD0274 merodiploid, we were able to show that ClosTron mutants could be obtained with high efficiency in this strain, but not in the parent $630\Delta erm$ wild type strain (Table 1). Previously, mutants made in this gene using single crossover plasmid integration developed no further than pin-prick colonies, and could not be propagated further [13]. Our data reinforces the view that this gene is essential and could represent a potential drug target for combating C. difficile infections. Interestingly, the equivalent gene in R20291 (CDR20291_0278 but misannotated as *metE*) was not designated as essential in the TraDIS study [7].

In this study, the necessary merodiploid cell was created through ACE-mediated inactivation of the *pyrE* locus. An alternative option would be to use a *pyrE* minus host and introduce the second copy of the gene into the genome concomitant with correction of the *pyrE* allele to wildtype. Restoration of uracil prototrophy (growth on minimal media without uracil supplementation) represents a simpler selective phenotype. More importantly, because the *pyrE* deletion mutation cannot revert, there can be no false positives. The required *pyrE* minus mutants of *C. difficile* strains R20291 and 630 are available [15], and are relatively easily created in any *Clostridium* [16].

In conclusion, we have developed a simple and rapid method that may be used to experimentally confirm essentiality in, but not limited to, *C. difficile*. Using this system, we demonstrated that *trpS*, *metK*, and *CD0274* are essential in *C. difficile* $630\Delta erm$. Our method may be used to confirm suggestions of essentiality that arise from genome-wide mutational methods such as TraDIS.

Authorship/contribution

DJFW undertook all the experimental work described in the study, with guidance from JTH, KW and NPM. JTH and NPM conceived the study. NPM and DJFW wrote the manuscript. All authors read and commented on the manuscript.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.anaerobe.2016.07.007.

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