



Tn6553, a Tn7-family transposon encoding putative iron uptake functions found in *Acinetobacter*

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Abstract

Acinetobacter baumannii is an opportunistic pathogen that has become difficult to eradicate mainly because of its high level of antibiotic resistance. Other features that contribute to this organism's success are the ability to compete for nutrients and iron. Recently, several novel Tn7-family transposons that encode synthesis and transport of siderophore and iron uptake systems were characterised. Here, another Tn7-type transposon (named Tn6553) is described. Tn6553 contains a set of iron utilisation genes with a transposition module related to Tn7. Tn7-family transposons that carry iron uptake systems facilitate the spread of these functions in *Acinetobacter* strains. Given that Tn7 is known to transpose efficiently into its preferred target site, finding siderophore functions on Tn7 family transposons is important in the context of dissemination of virulence genes amongst *Acinetobacter* strains.

Keywords *Acinetobacter baumannii* · Tn6553 · Tn7 family transposon · Iron acquisition

Introduction

Acinetobacter baumannii has proven to be a successful pathogen mainly due to high levels of antibiotic resistance, producing biofilm and the ability to compete for iron under iron-limiting conditions (Zimblet et al. 2009; Zarrilli et al. 2013; Harding et al. 2018). Their ability to compete for micronutrients required for growth, e.g. iron, contributes to their success allowing this organism to become a successful pathogen (Mortensen and Skaar 2012; Harding et al. 2018). Recently, we described a set of large novel transposons that carry genes for synthesising and transporting siderophores in several unrelated strains (Hamidian and Hall 2021). We showed that these transposons (Tn6171, Tn6552 and their variants) contain genes encoding transposition proteins related to those in Tn7 (Hamidian and Hall 2021) and that they can target the *glmS* gene, which is the preferred Tn7 target site (Craig 1991, 1996). Similar to Tn7 (Craig 1991,

1996), Tn6171, Tn6552 and their variants are flanked by five ($n = 5$) bp target site duplications (TSD) (Hamidian and Hall 2021). Here, the properties of yet another Tn7-family transposon designated Tn6553 that encodes iron uptake functions are described. This adds another mobile genetic element to the growing list of Tn7-family transposons that can disseminate important functions in *Acinetobacter* strains.

Methods

Sequence data used in this study

Complete genome sequences of the *Acinetobacter* strains, *Acinetobacter soli* strain GFJ2 and *A. baumannii* DS002 GenBank accession numbers CP027704.2 and CP016896, respectively, were used in this study to examine the structure of Tn6553 and its variant Tn6553-v1.

Sequence annotation and analysis

Standalone BLAST (available at <https://www.ncbi.nlm.nih.gov/books/NBK52640>) functions, including BLASTn, BLASTp and tBLASTn, were used for sequence analysis as previously described (Altschul et al. 1990). All coding regions of Tn6553 were found using the NCBI's Orf Finder

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program available at <https://www.ncbi.nlm.nih.gov/orffinder>. Gene features of transposons including all protein-coding regions were annotated manually using both Pfam (<http://pfam.xfam.org/>), UniProt (<https://www.uniprot.org>) and BLASTP (<http://blast.ncbi.nlm.nih.gov/Blast.cgi>) searches as described previously (Altschul et al. 1990; Finn et al. 2014, 2017). The IS-Finder database (<https://www-is.biotoul.fr/>) was used to identify insertion sequences (IS). Conserved sequences were determined using the WebLogo software available at <https://weblogo.berkeley.edu/logo.cgi>.

Results and discussion

A. baumannii strains have become successful opportunistic superbugs that have become difficult to treat because of the acquisition of genes conferring resistance to a wide range of antibiotics including carbapenems (Hamidian and Nigro 2019). Their success is also partly due to their ability to compete for micronutrients including iron (Mortensen and Skaar 2012; Harding et al. 2018). We previously described two Tn7-family transposons that carry iron uptake and siderophore functions that could increase the iron uptake capacity of strains that carry them by supplying the genes required for synthesis and uptake of the fimsbactin group of siderophores (Hamidian and Hall 2021). Here, searches of the complete genomes of *Acinetobacter* strains in GenBank non-redundant database using low stringency BLASTN and sequence of the boundaries of Tn6171 uncovered yet another Tn7 family transposon type, encoding a predicted iron uptake system. This transposon is located downstream of the chromosomal *glmS* gene of the *Acinetobacter soli* strain GFJ2 (GenBank accession number CP016896) and is flanked by five ($n=5$) bp TSD (Fig. 1a, b). This transposon is novel and hence was herein named Tn6553. Tn6553 is a 23791 bp Tn7 family transposon, bounded by 28 bp imperfect inverted repeats, and like Tn6171 and Tn6552, further transposon binding sites for TnsB were detected within 200 bp of the Tn boundaries (Fig. 1b, c). Tn6553 includes a complete set of genes (*tnsABCDE*) encoding transposition proteins with 41–48% aa identities to TnsABCD and 26% identity to TnsE encoded by Tn7 (and with 32–43% aa compared to TnsABCDE in Tn6171). However, the *tns* module is interrupted by the iron acquisition segment located between the *tnsD* and *tnsE* genes.

Tn6553 encodes a set of functions required for iron uptake. It contains several genes such as *irtAB*, *bauDCEBA* and *fur*, which mainly encode proteins predicted to be involved in ferrichrome uptake and transmembrane transport (Fig. 1a), which are distantly related to *Escherichia coli* ferrichrome-iron receptor (Coulton et al. 1983). The *irtAB* genes encode putative ABC-type multidrug transport systems (ATPase and permease component)

belonging to the MdiB superfamily (Protein family accession number COG1132). The *bauDCE* genes encode putative iron chelate uptake ABC transporter family permease subunit (ferric acinetobactin ABC transporter), while *bauB* predicted to encode a siderophore-binding periplasmic lipoprotein and *bauA* a TonB-dependent siderophore protein (Fig. 1a). Tn6553 also carries a *fur* gene encoding an Fe²⁺ (or Zn²⁺) uptake transcriptional regulation protein (Protein family accession number COG0732), which is in fact a putative master transcriptional regulator of the iron-responsive genes. Notably, the iron uptake proteins encoded by Tn6553 are distantly related to the siderophore systems (with amino acid identities ranging from 20–30%) that we recently described in *A. baumannii* in different Tn7-family transposons (Hamidian and Hall 2021). However, whether this operon functions as a siderophore system remains to be experimentally examined.

A recent study reported that *A. baumannii* DS002 (GenBank accession number CP027704.2) contains a novel siderophore system present in a genomic island that encodes 30 genes/orfs including four ($n=4$) genes encoding putative transposases TnsBC1C2E, 13 gene encoding putative iron acquisition functions, and the 13 reading frames coding for proteins of unknown functions (Yakkala et al. 2019). However, detailed sequence analysis showed that this iron uptake system is in a variant of Tn6553, here called Tn6553-v1 (Fig. 1a). Tn6553-v1 is 24,839 bp in size. It encodes 23 open reading frames, including transposition functions, iron acquisition, DNA metabolism and four ($n=4$) hypothetical proteins. Tn6553-v1 is 1047 bp longer in size than Tn6553 due to the insertion of a novel 1039 bp IS5 family insertion sequence, named ISAbA53 (Fig. 1a), and its eight ($n=8$) bp TSD. In addition to the presence of a copy of ISAbA53 (a novel IS5 family insertion sequence that is 85% identical to ISAha1) in Tn6553-v1, it also differed from Tn6553 by 17 bp and six ($n=6$) bp gaps across the remaining segments. Notably, the *tnsC* gene is split into two ($n=2$) reading frames (Fig. 1a), likely due to sequencing errors. The *itrB* gene was also divided into two ($n=2$) frames, probably due to sequencing or assembly errors. However, these could not be verified here as the strain was not accessible. Together, the Tn7 family transposon found in DS002 (Tn6553-v1) is an ISAbA53 interrupted variant of Tn6553 (Tn6553::ISAbA53), indicating that Tn6553 is more ancestral and that *A. soli* might be the source. This is consistent with both isolates recovered and evolved in the same environmental niche, soil. Differences in TniABCDE of Tn6553 compared to those in Tn7, Tn6171 and Tn6552 (~48–52% aa identity) indicate that these Tn7 family transposons belong to a very diverse family.

Searching the GenBank non-redundant database; however, a region distantly related to the putative iron uptake region of Tn6553 (with approx. 66–69% DNA identity)

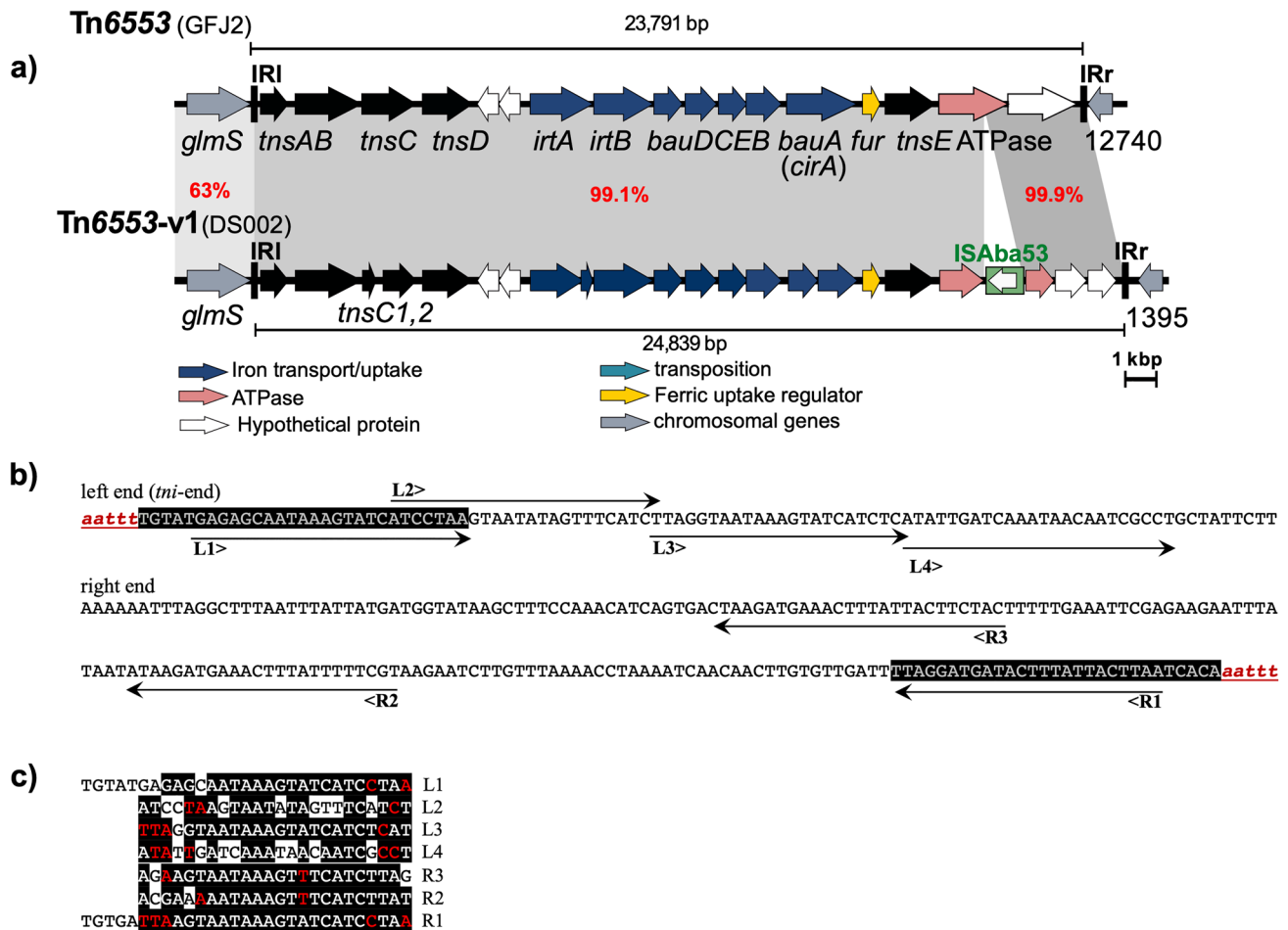


Fig. 1 Genetic structures of Tn6553 and Tn6553-v1 **a**, Inverted repeats (IR) of Tn6553 **b** and sequence conservation in transposon binding sites of Tn6553 **c**. In **a**, horizontal arrows indicate the direction and orientation of genes. Transposition genes are coloured black, iron uptake systems are dark blue, chromosomal genes are grey and white indicating hypothetical proteins. **b**, IR and transposon binding sites of Tn6553. Target site duplications are in lower case and red

(5 bp). The left and right end of Tn6553 is shown using uppercase letters with IRs grey on black. Horizontal arrows below and above the sequence lines (L1-4 and R1-3) are transposon binding sites, TBSs. **c**, sequence conservation of IRs and TBSs. Conserved sequences are shown using white or red letters on black (on the left) and a WebLogo (on the right) generated using the software available at <https://weblogo.berkeley.edu/logo.cgi>

was found in several *Acinetobacter* strains, namely *Acinetobacter* sp. Tol 5 DNA, *Acinetobacter bereziniae* strain

GD03255, *Acinetobacter bereziniae* strain GD03393 (shown as a representative in Fig. 2), *A. bereziniae* strain GD03185,

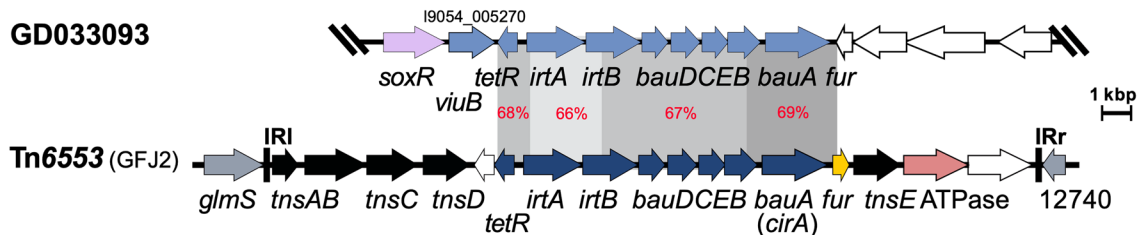


Fig. 2 Comparison of the putative iron uptake region of Tn6553 with the corresponding region in *Acinetobacter bereziniae* strain GD03393. Horizontal arrows indicate the direction and orientation of genes. Arrows coloured different shades of blue represent putative iron uptake genes. Transposition genes of Tn6553 are coloured black.

Scale bar is shown. Regions with significant homology are shown using shades of grey with their DNA identities indicates in red. Figure is drawn to scale from GenBank accession numbers CP092085 (GD03393 chromosome) and CP016896 (GFJ2 chromosome)

A. bereziniae strain XH901 and *Acinetobacter guillouiae* NBRC 110,550 (GenBank accession numbers, CP092085 AP024708.1, CP092083.1, CP066119.1, CP018259.1 and AP014630.1) (Fig. 2). This included an approximately 8 kb segment starting from the 5'-end of the *tetR* gene extending to ~100 bp downstream of the *bauA* gene (Fig. 2). Notably, this putative iron uptake region included an additional gene, namely *viuB* (locus id_19054_005270 in GenBank accession number CP092085), encoding a NADPH-dependent ferric siderophore reductase. This gene is missing in Tn6553. These *Acinetobacter* strains belong to different species (*berezinae* and *guillouiae*) and given that they are recovered from diverse clinical and environmental samples in different geographical regions indicating that this putative iron uptake system is not restricted to a specific species, region, or niche. However, given that no putative iron uptake region with significant homology was found outside the *Acinetobacter* genus also suggests that *Acinetobacter* is the origin.

This work describes yet another novel Tn7 family transposon type that encodes a predicted iron uptake system. Despite evolutionary differences these transposons share the same properties and are found in the same chromosomal position as Tn7. Finding siderophore functions on Tn7 family transposons in an important finding in the context of dissemination of virulence genes amongst *Acinetobacter* strains given that Tn7 is known to transpose very efficiently into its preferred target site. The presence of siderophore/iron uptake systems on Tn7-family transposons facilitates the spread of these functions providing further evidence on strategies that *Acinetobacter* strains use to rapidly evolve, compete, and adapt under extreme conditions. None of the transposons that carry iron uptake/siderophore function characterised to date (Tn6171, Tn6552 and Tn6553) have been found outside the *Acinetobacter* genus, however, as more genomes are sequenced additional related transposon might likely be found in this genus and maybe in other bacterial genera.

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Availability of data and materials All genome sequence data analysed during this study are publicly available in GenBank and cited in the article.

Declarations

Competing interests The authors declare no competing interests.

Ethics approval and consent Not applicable.

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