Genomic Analysis of Molecular Bacterial Mechanisms of Resistance to Phage Infection

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Abstract

Background

In order to optimize phage therapy, we need to understand how bacteria evolve against phage attack. One of the main problems of the phage therapy is the appearance of bacterial resistance variants. The use of genomics to track antimicrobial resistance is increasingly developed and used in clinical laboratories. For that reason, it is important to consider, in an emerging future with phage therapy, to detect and avoid phage resistant strains, that can be overcome by the analysis of metadata provided by WGS. Here, we identified genes associated with phage resistance in 18 *Acinetobacter baumannii* clinical strain belonging to the ST-2 clonal complex during a decade (Ab2000 vs 2010): 9 from 2000 and 9 from 2010.

Results

The presence of genes putatively associated to phage resistance were detected. Genes detected were associated with an abortive infection system, restriction-modification system, genes predicted to be associated with defence systems but with unknown function and CRISPR-Cas system. Between 118 and 171 genes were found in the 18 clinical strains. On average, 26% of these genes were detected inside genomic islands (GIs) in the 2000 strains and 32% in 2010 strains. Furthermore, 38 potential CRISPR arrays in 17 of 18 of the strains were found, as well as 705 proteins associated with CRISPR-Cas systems.

Conclusions

A moderately higher presence of these genes in the strains of the 2010 in comparison to those of the 2000 were found, especially those related to the R-M system and CRISPR-Cas system. The presence of these genes in GIs in a higher rate in the strains of the 2010 compared to those of the 2000 was also detected. WGS and bioinformatics could be powerful tools to avoid drawbacks when a personalized therapy is applied. In this study, it allows us to take care of the phage resistance in *A. baumannii* clinical strains to prevent a failure in a possible phage therapy.

Background

As part of the ESKAPE pathogens, *A. baumannii* is frequently isolated from infections in clinical environments, and its resistance against multiple antibiotics is increasingly common (1). For this reason, it is necessary to opt for alternative treatments, such as phage therapy. However, the ability of bacteria to develop resistance mechanisms against phages is possible, even when there is no previous treatment with phage therapy due to the constant coevolutionary interactions (2). The spread of phage resistance presents a significant challenge to the efficacy of the therapy (3), (4).

It is important to know and characterize the phage resistance mechanisms of a certain species, clone or strain, prior to phage treatment in order to minimise treatment failure. Whole-genome sequencing (WGS) has been demonstrated to be a powerful tool in the detection of phage-resistance mechanisms, as well as the evolution of CRISPR-Cas arrays in bacteria subjected to phage pressure (5, 6). WGS is increasingly becoming
a cheaper and faster technology, thus it is implemented progressively in routine hospital diagnostics and research (7).

Recently, new or modified phage resistance mechanisms have been discovered and characterized (8). Although a large part of defence systems against phages are maintained over generations, there is a continuous emergence of resistance mechanisms due to spontaneous mutations as a consequence of the coexistence of phage and bacteria. Most of these mutations occur in the phage receptors proteins, employed by the phages to adhere to the cell (8). In recent years, phage resistance mechanisms are attracting increasing interest due to the rising knowledge in phage interactions with bacteria. This is leading to the discovery and characterization of new phage resistance mechanisms such as Zorya, Druantia or Thoeris (9). Phage resistance mechanisms are typically clustered in genomic “defence islands”: mobile genetic evolutionary elements that contain genes associated with phage defence systems (9, 10).

The main resistance mechanisms are related to the inhibition of the phage adsorption, blocking of phage DNA injection, cutting of the injected DNA, inhibition of the phage DNA replication, interference in the phage assembly, and bacterial suicide (11). In Fig. 1 we summarized all the characterized phage resistance mechanisms.

In this study, we focused in those which could be bioinformatically detected without any experimental process:

i) Abortive Infection systems, characterized by the fact that the phage enters the cell, but its development is interrupted in any phase (replication, transcription or translation). The mechanism of action is not entirely clear, either because of their complexity or because they are widely varied from one species to another (12).

ii) Toxin/Antitoxin systems are a specific type of ABI system but they are well-characterized and widespread through diverse species (13). In this system, a toxin is produced by the cell and is neutralized by an antitoxin. The expression of these molecules is highly controlled and varies from one system to another. When the balance between one molecule and the other is disturbed, the toxin is released and the bacteria die (14).

iii) Restriction-Modification systems consist of a restriction endonuclease and a This type of system distinguishes the DNA of the host from foreign DNA to recognize and destroy phage DNA after its injection into the When unmethylated phage DNA enters a bacteria which possesses the R-M system, it will be cleaved by the restriction endonuclease or methylated by the methyltransferase to escape the restriction (15).

iv) CRISPR-Cas (clustered regularly interspaced short palindromic repeats – CRISPR-associated system is an adaptative immune system that bacteria develop against phage DNA/RNA and other foreign DNA (16). The typical structure of the CRISPR-Cas locus is a leader sequence, followed by the repeat-spacer array and the cas genes operon (17). The adaptation of the CRISPR-Cas system is due to the acquisition of the spacer sequences, which are small fragments of foreign nucleic acids, between the repeats of the CRISPR locus (17). The functioning of the CRISPR-Cas system, usually divided into three steps (adaptation, processing and guidance of the crRNA-CRISPR RNA- and targeting and interference of the foreign DNA/RNA) is carried out by the Cas (CRISPR-associated) proteins (18). CRISPR-Cas systems are classified according their conserved cas genes and the architecture of the cas operon (19). Until recently, little data existed about CRISPR-Cas systems in A. baumannii. The pangenome analysis of A. baumannii has shown CRISPR-Cas
systems in the species (20). One of the most characterized systems in *A. baumannii* is the CRISPR I-Fb system (21). However, most of the Cas-related genes and CRISPR arrays are yet not identified and characterized.

In this study, we searched for putative genes associated with phage resistance and we focused on CRISPR-Cas systems by studying the CRISPR arrays and Cas protein presence through bioinformatic approach in 18 genomes of clinical strains of *A. baumannii* isolated in the “II Spanish Study of *A. baumannii* GEIH-REIPI 2000–2010”.

**Methodology**

**Genome database**

18 clinical *A. baumannii* genomes previously sequenced and annotated (II Spanish Multicenter Study, GEIH-REIPI Acinetobacter baumannii 2000–2010; Umbrella Bioproject PRJNA422585) (22), (23) have been studied. Nine strains were from the 2000 and nine from 2010. All of the strains belong to the ST-2 clone (22).

**Search for general genes associated to bacteriophage resistance and their presence in genomic islands**

In order to analyse the presence of genes putatively associated with phage resistance systems, a custom database based on genes from the “PADS Arsenal database” (https://bigd.big.ac.cn/padsarsenal/) was created (24). The genes were grouped in five systems: ABI systems related (not belonging to toxin/antitoxin system), TA systems related, R-M system related, CRISPR-Cas associated proteins and newly (NEW) characterized systems related genes. In this last category we included those genes which hit against known phage resistance genes but were associated with genes predicted to be associated with phage resistance functions and whose function in *A. baumannii* is not clear yet, such as newly characterized systems (e.g. Zorya, Druantia, Thoeris). A blast search of the complete genomes against this database and filtered out those hits which e-value > 1E-04 was made. The percentage of the genes involved in resistance was calculated by dividing the genes predicted to be associated with phage resistance by the total number of genes in the bacteria genome.

To locate Genomic Islands (GIs) three different approaches were used: IslandViewer with default settings, (25), blast search with default settings and cut-off of e-value < 1e-03 against a previously constructed ICEberg database (26) and checking the Guanine-Cytosine (GC) content of the contigs of each genome (27). The previously detected phage-resistance genes were localised in the GIs detected per genome and average percentage of genes by collection was calculated.

**Search and characterization of CRISPR arrays**

In a first try, CRISPRCasFinder (28) was employed but no putative CRISPR-Cas system was found. For this reason, CRISPR arrays were found using the CRISPR Recognition Tool (CRT) (29). The modification proposed by Rho et al (29) for whole-metagenomic assembled genomes called metaCRT was used (30) with
the following parameters: minimum number of repeats: 3, minimum repeat length: 12, maximum repeat length: 70, minimum spacer length: 18, maximum spacer length: 80 and with a search window of: 6.

In order to filter and validate the CRISPR arrays, a similar procedure to the first step in the protocol developed by Shmakov et al was followed (31) (Fig. 2). In the first step, CRISPR arrays separated by no more than 6 Open Reading Frames (ORF) to a putative Cas protein identified before, were considered to be part of a putative CRISPR-Cas system. Those which not were part of a putative CRISPR-Cas system were considered to be single possible CRISPR arrays. To validate these arrays, we made a short-blastn search of the spacers of the possible CRISPR arrays against all phage genomes using the INPHARED database (32). Results that were > 95% in identity and those arrays whose query hit was larger than 20 were considered putative CRISPR arrays.

Negative blast hits of the spacers of the CRISPR arrays against bacteriophage follow the procedure described by Shmakov for isolated arrays: first, arrays > 400 bp and with an ORF coverage > 0.95 were filtered out. Second, all arrays with < 850 bp that had domain in the Conserved Domain Database (CDD) search were filtered out (33), (34). Then, pairwise distances between spacers of each array were calculated (number of matches in the longest blastn hit between them, divided by the length of the smaller spacer in the pair). The spacers of each array were clustered using single linkage clustering following the same procedure as Shmakov et al. with a cut-off of 0.3. A spacer similarity index was calculated for each CRISPR array as the number of clusters formed divided by the number of spacers in the array (1 means that all the spacers are different). Those arrays whose spacer similarity index was < 0.85 were filtered out. The rest were considered putative CRISPR arrays.

To complement the procedure made by Shmakov, a search about common special low-complexity sequences that may be confused as CRISPR arrays was made, known as false-CRISPR elements (35). The presence of Tandem repeats, potentially hypermutable regions which enable bacteria to adapt to evolving environments without increasing their mutation rate, was checked with Tandem Repeat Finder (36–38). The presence of short low-complexity repeats was also examined with RepeatMasker (39). To test the results and to complete the search of low-complexity sequences a blast search against an existing false-CRISPR elements database was obtained from the CRISPRone website (35).

The search for possible Cas-related proteins was made, based on the method of Zhang et al. (40), but adding a search in "HMMCAS" website of all of their available HMM models, performed with a cut-off of reported e-values of (41). The 18 genomes were examined using hmmscan with all of the pfam HMM profiles based on NCBI entries of known Cas protein families searching in the pfam database 70 Cas-related protein families and other CRISPR-associated proteins in the pfam database (e. g. DEAD/DEAH box helicase), 93 families which were in the TIGRFAMS resource and the 24 newly characterized families (40, 42–44). For the TIGRFAMS database proteins it was necessary to build an HMM profile with hmmbuild with default settings (42) making a previous alignment for each compound of proteins with Clustal Omega version 1.2.4 (ClustalO) (45).

All of the input and output data for the searches (genes associated to phage resistance, genomic islands, CRISPR arrays and Cas proteins) were processed with Python (www.python.org) and BioPython (19304878
In order to establish the evolution and to compare the presence of the CRISPR arrays among the same clonal complex of the 18 clinical strains of *A. baumannii*, a phylogenetic tree was made using the CRISPR spacers detected. Trees were built using MEGA7 with CLUSTALW alignment (MEGA version 7.9.26) (48–50).

**Results**

**Genes putatively associated with phage resistance in A. baumannii clinical strains and their presence in GIs**

Between 118 and 171 genes were detected per genome, those could be putatively associated with bacterial defence against bacteriophages (Table 1, Additional file 1). The frequency (%) of each resistance system was calculated with the number of genes of each group divided by the total genes per genome. It was observed that the genes related to R-M and CRISPR-Cas systems showed a slightly higher prevalence in 2010 strains (Fig. 3, A). The frequency of the genes related to ABI, TA and new systems remained constant in both collections. The presence of putative phage resistance genes in GIs was also predicted (Table 2, Additional file 1), and it was found that in GIs represents in strains of the year 2010 approximately a 24% in average and approximately a 19% in the strains of the year 2000 in average (Fig. 3, B). The observed increase was produced specially in genes related to the RM system, to those related with new phage resistance mechanisms and CRISPR-Cas system.

**CRISPR arrays**

180 putative predicted arrays were found (without filtering) using metaCRT in the 18 genomes of ST-2 *A. baumannii* clinical strains (Table 3, Additional file 2) Post the complete process of filtering designed by Shmakov et al. (31) and removing the low-complexity sequences (35), only 40 CRISPR arrays were selected (Table 1): 18 CRISPR arrays were present in the 2000 strains and 22 in the 2010 strains. All the strains, excepting in the Ab161_GEIH-2000 strain, presented at least 1 CRISPR array.
Table 1
CRISPR arrays present in the genomes of 18 A. baumannii clinical strains.

| Strain          | Contig         | Size | Start | Stop  | Repeat                  | Nº of spacers |
|-----------------|----------------|------|-------|-------|-------------------------|---------------|
| Ab33_GEIH-2010  | MSMK01000003   | 160  | 10462 | 10622 | ATTTTGAATTTAAAAA        | 4             |
| Ab33_GEIH-2010  | MSMK01000187   | 198  | 18280 | 18478 | ACAAAAGAAAAAT           | 4             |
| Ab49_GEIH-2010  | MSMM01000317   | 96   | 1114  | 1210  | TCATTTTGCTGTTGTT        | 2             |
| Ab49_GEIH-2010  | MSMM01000323   | 198  | 78    | 276   | ACAAAAGAAAAAT           | 4             |
| Ab49_GEIH-2010  | MSMM01000347   | 122  | 367   | 489   | TTTAAATTCAAAA           | 3             |
| Ab54_GEIH-2010  | MSML01000240   | 198  | 3914  | 4112  | AATTTTTTTTCT            | 4             |
| Ab54_GEIH-2010  | MSML01000469   | 96   | 1108  | 1204  | TCATTTTGCTGTTGTT        | 2             |
| Ab54_GEIH-2010  | MSML01000525   | 164  | 8017  | 8181  | ATATATTTTGAA           | 3             |
| Ab76_GEIH-2010  | MSLY01000008   | 96   | 835   | 931   | TCATTTTGCTGTTGTT        | 2             |
| Ab76_GEIH-2010  | MSLY01000677   | 198  | 3369  | 3567  | AATTTTTTTTCT            | 4             |
| Ab76_GEIH-2010  | MSLY01000708   | 164  | 714   | 878   | ATATATTTTGAA           | 3             |
| Ab103_GEIH-2010 | MSLX01000655   | 160  | 9148  | 9308  | ATTTTGAATTTAAAA         | 4             |
| Ab103_GEIH-2010 | MSLX01000266   | 164  | 2680  | 2844  | ATATATTTTGAA           | 3             |
| Ab103_GEIH-2010 | MSLX01000388   | 96   | 1108  | 1204  | TCATTTTGCTGTTGTT        | 2             |
| Ab103_GEIH-2010 | MSLX01000506   | 198  | 55    | 253   | ACAAAAGAAAAAT           | 4             |
| Ab104_GEIH-2010 | MSMA01000019   | 96   | 1450  | 1546  | TCATTTTGCTGTTGTT        | 2             |
| Ab104_GEIH-2010 | MSMA01000107   | 160  | 4402  | 4562  | TTTAAATTCAAAA          | 4             |
| Ab104_GEIH-2010 | MSMA01000246   | 164  | 10815 | 10979 | ATATATTTTGAA           | 3             |
| Strain         | Contig       | Size | Start  | Stop   | Repeat                   | Nº of spacers | spacers |
|---------------|--------------|------|--------|--------|--------------------------|---------------|---------|
| Ab105_GEIH-2010 | LJHB01000001 | 198  | 125508 | 125706 | ACAAAAAGAAAAAT           | 4             |         |
| Ab105_GEIH-2010 | LJHB01000010 | 292  | 7321   | 7613   | TAAAATAATTTTAA           | 5             |         |
| Ab121_GEIH-2010 | MSLZ01000141 | 198  | 4992   | 5190   | AATTTTTCTTTTCT           | 4             |         |
| Ab122_GEIH-2010 | MSMD01000782 | 164  | 711    | 875    | ATATATTTTTGA            | 3             |         |
| Ab155_GEIH-2000 | LJHA01000001 | 198  | 125512 | 125710 | ACAAAAAGAAAAAT           | 4             |         |
| Ab155_GEIH-2000 | LJHA01000002 | 292  | 7323   | 7615   | TAAAATAATTTTAA           | 5             |         |
| Ab158_GEIH-2000 | MSMC01000196 | 198  | 4027   | 4225   | AATTTTTCTTTTCT           | 4             |         |
| Ab158_GEIH-2000 | MSMC01000525 | 136  | 868    | 1004   | ATTTTTAATATTTA          | 3             |         |
| Ab166_GEIH-2000 | MSMG01000383 | 86   | 859    | 945    | AAATAGCCTAAGC           | 2             |         |
| Ab166_GEIH-2000 | MSMG01001001 | 198  | 293    | 491    | ACAAAAAGAAAAAT           | 4             |         |
| Ab166_GEIH-2000 | MSMG01000974 | 79   | 1310   | 1389   | TCTGCTGTCGAAAA           | 2             |         |
| Ab166_GEIH-2000 | MSMG01001128 | 194  | 304    | 498    | ACGACGTGGACGATCTTC       | 3             |         |
| Ab169_GEIH-2000 | MSMF01000039 | 96   | 797    | 893    | TCATTTTGCTGTGGTT         | 2             |         |
| Ab169_GEIH-2000 | MSMF01000336 | 198  | 152    | 350    | ACAAAAAGAAAAAT           | 4             |         |
| Ab175_GEIH-2000 | MSMI01000153 | 79   | 8115   | 8194   | TTTCCGACAGCAGA          | 2             |         |
| Ab175_GEIH-2000 | MSMI01000682 | 86   | 2355   | 2441   | AAATAGCCTAAGC           | 2             |         |
| Ab177_GEIH-2000 | MSME01000459 | 198  | 215    | 413    | ACAAAAAGAAAAAT           | 4             |         |
| Ab183_GEIH-2000 | MSMJ01000620 | 96   | 1077   | 1173   | TCATTTTGCTGTGGTT         | 2             |         |
| Ab183_GEIH-2000 | MSMJ01000380 | 198  | 78     | 276    | ACAAAAAGAAAAAT           | 4             |         |
| Strain           | Contig          | Size | Start | Stop  | Repeat                      | Nº of spacers |
|------------------|-----------------|------|-------|-------|-----------------------------|---------------|
| Ab192_GEIH-2000  | MSMH01000263    | 96   | 1139  | 1235  | TCATTTTGCTGTTGTT            | 2             |
| Ab192_GEIH-2000  | MSMH01000273    | 157  | 0     | 157   | TTGAATTAAAAA                | 4             |
| Ab192_GEIH-2000  | MSMH01000395    | 198  | 21634 | 21832 | AAAAAAGAAAAAT              | 4             |

A phylogenetic tree of the complete CRISPR array sequences was constructed (Fig. 4), and showed an equal distribution of the spacers between the strains of the two years. Some of the spacers were predicted to be the same even in strains different year collections. Few of the arrays were unique respect to the other, such as the present in the 2000 strains Ab158_GEIH-2000_MSMC01000525, Ab166_MSMG01000383, Ab166_MSMG01000974, Ab166_MSMG01001128, Ab175_MSMI01000153 or Ab175_ MSMI01000682. However, there were 5 CRISPR arrays grouped that only were represented in the 2010 strains (Fig. 4).

**Cas-related proteins**

When HMM against Cas-known, Cas-related and CRISPR-associated protein families was employed, 705 Cas-related proteins were identified in the 18 genomes: 341 Cas-related proteins were detected in 2000 strains and 364 in 2010 strains (Table 4, Additional file 3). Most of them were identified as DEAD/DEAH box helicase (207 of the total) and as Type III Restriction Unit Res III (195 of 705). The vast majority of them were located next to proteins whose predicted function does not match with a Cas protein function or to proteins whose function was unknown. Other Cas-related were close in the same contig thereby giving us a clue to help identifying a functional Cas cluster. For example, in the contig MSLX01000260 from the Ab103_GEIH-2010 strain a putative Helicase_C protein (OLV37994.1) and a Cas_St_Csn2 protein (OLV37998.1) were only of 2 ORF distance between them. However, the function of the surrounding proteins was hypothetical, thus hindering the identification process as a Cas cluster.

**Discussion**

In clinical laboratories, genomics is rapidly being developed and utilized to track antibiotic resistance. As a result, it is critical to explore how to detect and avoid phage resistant strains, if a treatment based on phages was going to be applied, by using WGS metadata analysis. In this study, we looked for genes linked to phage resistance in 18 clinical strains of *A. baumannii*. We constructed a database with genes based in the public PADS database, as it is the most complete database about prokaryotic antiviral defence systems so far, as well as being collecting newly discovered types of defence systems to the BIG Data Center (24, 51). In this case, the high number of genes made us establish groups in order to simplify the results of the blast hits. We also tried to identify the presence of CRISPR-Cas systems by separating the search in CRISPR arrays and Cas proteins.

A difference between the presence of phage resistance genes in 2010 strains and 2000 strains was observed, with a higher presence of genes related to the RM system and CRISPR-Cas system and lower of
TA-related genes. The natural reciprocal selection pressure between host bacteria and phage increases the infectivity of the phage and the phage-resistance in the bacterium side (52). In fact, phage populations are ubiquitous at body surfaces such as lungs, intestines or skin, and they outnumber bacteria at least by 10-fold (53). In this study, the acquisition of phage resistance genes is correlated with a higher presence of complete prophages in the strains of the 2010 in comparison with those of 2000 (54). This could be a result of the development of phage resistance adaptative systems, that could promote the emergence of new phages that can overcome them, such as could happen with Ab105-1ϕ and Ab105-2ϕ, two prophages present in the 2010 collection strains but not in the 2000 collection strains (54).

Defence systems are regularly obtained by bacteria and archaea through horizontal gene transfer (HGT) owing to environmental adaptation of the bacterial communities (55, 56). We found a major average of genes acquired by HGT in the 2010 strains rather than in the 2000 ones, especially those genes related to RM system and CRISPR-Cas. It was demonstrated that only ~ 4% of RM systems are in the core genomes of prokaryotic species, suggesting they are commonly transferred (57). CRISPR-Cas systems display weak consistency within the core genome, demonstrating the prevalence of the HGT spreading this system (57, 58). The RM system and the CRISPR-Cas system commonly coexist with an elevated contribution to the bacterial immunity and they rarely operate on their own (57, 59). However, they are far from being perfect in the bacterial resistance, and phage can escape these systems by many different ways, for example the anti-CRISPR proteins (8, 60). We also observed a decreasing number of TA-related genes through the years, even their presence in GIs is higher in the 2000 strains than in the 2010’s. This could be because the counteradaptation of the phage may be reached by developing antitoxin in the phage genome that inhibit the cell death and thus promote the infection of the phage (61, 62) or because they could have evolved into Cas proteins of the CRISPR-Cas system, as the TA proteins are considered as ancestors of Cas2 proteins (63).

Furthermore, we found the CRISPR-Cas genes blast hit results incomplete due to the separation in contig assembly of the genomes, which prevented us from identify proteins or arrays related to the CRISPR-Cas proteins identified in small contigs (data not shown), and also due to the high diversity of the Cas proteins and the little knowledge about these proteins in clinical strains of A. baumannii, which increases the difficulty identifying these type of proteins (20, 64). As a consequence, we examined the presence of CRISPR arrays and Cas proteins separately. We establish a methodology to discard false-CRISPR elements based on the method of Shmakov et al. (65) and posteriorly completed with a full evaluation of the quality of the CRISPR arrays filtered based on the search of tandem repeats, simple repeats and their presence on phage genomes (35). Secondly, another reason of developing an alternative method is the nature of the multi-resistant pathogens, their constant adaptation to different environments and thus the continuous acquisition of different mobile elements, which provokes the appearance of new CRISPR-Cas yet to be identified (66). This also fosters and extends the variability in the Cas proteins, complicating their characterization.

40 CRISPR arrays were found in the 18 A. baumannii clinical strains from the ST-2 clone. All of the strains presented at least one CRISPR array except one, Ab161_GEIH-2000. The vast majority of the arrays are shared between the clone ST-2 in both collections, with some exceptions such as the five arrays only found
in 2010 strains. It has been shown that the distribution of CRISPR-Cas system is MLST dependent and non-random, and thought to be a better discriminating tool than classical MLST in discriminating different *K. pneumoniae* (67, 68). On the other hand, the detection of different unique CRISPR arrays only in the 2000 strains demonstrates the dynamic interaction of these arrays throughout the years.

All of the CRISPR arrays in this study were without any Cas or putative Cas protein near to them. It was described that these “orphan” arrays belong to unknown CRISPR-Cas systems due to be to an extremely evolutionarily remote type of CRISPR-Cas (65). This existence of isolated CRISPR arrays could be explained for four reasons. First, the contig format of the studied genomes could provoke that some arrays are detected in small or incomplete contigs. Secondly, some Cas endonucleases such as Cas1 and/or Cas6 can recognize remote CRISPR arrays (69, 70). Third, it may occur the possibility of some of the unique isolated arrays form part of an undescribed CRISPR-Cas cluster extremely distant to the ones already characterized (65). And fourth, the strains may have lost the *cas* genes thus leaving the isolated arrays (65). The Cas distribution observed in this work would correspond and complete any of the hypothesis about the explanation of “orphan” CRISPR arrays mentioned before as the putative Cas proteins hit through the HMM search could form part of a complete Cas cluster. However, as it was said at the ending of the results section, it was impossible to determine *in silico* if the putative Cas detected form part of a complete and functional Cas loci.

The localization and characterization of defence systems against phages is a necessary step when designing an effective phage therapy. The WGS combined with an effective bioinformatics strategy would allow us to know what mechanisms the clinical strains have. This study shows the wide presence of genes associated with resistance against phages and their acquisition by GIs for 10 years in clinical *A. baumannii* strains from the same clonal complex ST-2 and the CRISPR arrays present on them.

**Abbreviations**

WGS: Whole Genome Sequencing; ST: Sequence Type; CRISPR: Clustered Regularly Interspaced Short Palindromic Repeats; GI: Genomic Island; ESKAPE: *Enterococcus faecium*, *Staphylococcus aureus*, *Klebsiella pneumoniae*, *Acinetobacter baumannii*, *Pseudomonas aeruginosa* and *Enterobacter spp.*; ABI: ABortive Infection; TA: Toxin Antitoxin; R-M: Restriction-Modification; GEIH: Grupo de Estudio de Infección Hospitalaria (Hospitalary Infection Group of Study); GC: Guanine Cytosine; CRT: CRISPR Recognition Tool; ORF: Open Reading Frames; CDD: Conserved Domain Database; HMM: Hidden Markov Model; PADS: Prokaryotic Antiviral Defence System; HGT: Horizontal Gene Transfer; MLST: Multi-Locus Sequence Typing

**Declarations**

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**Author contributions**

A.A., L.B., M.L., O.P. L.F-G., I.B., developed the analysis of results and wrote manuscript. C.O.C., A.M., revised manuscript. M.T., financed and directed the experiments as well as supervised the writing of the manuscript.

**Transparency declarations**

The authors have not actions to declare

**Availability of data and materials**

The datasets analysed during the current study are genomes of the "II Spanish Multicenter Study. GEIH-REIPI Acinetobacter baumannii 2000-2010" available in the BioProject PRJNA422585 repository with the link [https://www.ncbi.nlm.nih.gov/bioproject/422585](https://www.ncbi.nlm.nih.gov/bioproject/422585)

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Figures
Figure 1

Representation of the main mechanisms of bacterial resistance against phage infection (from the left in a clockwise sense). 1. a: The phage recognizes the bacterial membrane receptor and can carry out the infection; b: Alterations in the receptors are produced by mutations and prevent the phage from recognizing the receptor, so it will not infect the bacteria; c: The bacteria can block recognition by producing inhibitors that bind to receptors. 2. Production of exopolysaccharide or extracellular matrix. 3. OMVs are composed of membrane lipids, membrane proteins and periplasmic components and they are as a decoy against phages as a defence mechanism. 4. a. Bacteria block the injection of DNA from other phages, acquiring Sie systems through prophages with this type of protein; b. Once the bacterium has the prophage in its genome with the proteins that code for the Sie system, it will be able to block the entry of DNA from other phages. 5. a. The R-M system distinguishes between methylated and unmethylated DNA. Restriction enzymes cannot cut methylated DNA; b: If the phage DNA is not methylated, this system can cut the injected DNA. 6. CRISPR-Cas recognizing phage DNA sequences, incorporating them into the system and producing enzymes that are capable of recognizing these sequences to cut them. 7. The PICI system is found in the bacterial genome and induced by helper prophages to produce mature phage particles that assemble the PICI system itself to kill the infected cell and spreading this system to adjacent cells. 8. a. The most characterized Abi system is the toxin-antitoxin system. Under normal conditions, the bacterium expresses both proteins equally, so cell death does not occur; b. When the organism is subjected to stress situations, such as phage infection, the toxin is highly expressed in comparison with the antitoxin, causing cell death.
Figure 2

Search procedure of CRISPR-Cas systems in 18 genomes of A. baumannii clinical strains. Orange rectangle represents bioinformatic programmes used for that task. Green plus marks represent a positive result for each operation. Red minus marks represent a negative result for each operation.
Figure 3

A. Frequency (% rounded to two decimal numbers) of each group of genes in each genome respect to the total of genes: ABI (Abortive Infection System), TA (Toxin/Antitoxin system), RM (Restriction-Modification system) and NEW (genes associated with newly phage resistance bacterial mechanisms, e.g., Zorya, Hachiman, Druantia). B. Presence and non-presence of the putative phage resistance genes in GIs. The presence (%) rounded without decimal numbers section is divided into the different groups of genes.
**Figure 4**

Phylogenetic classification of the CRISPR arrays detected in 18 genomes of *A. baumannii* ST-2 clinical strains through a maximum-likelihood tree with the suggested model by the MEGA analysis Tamura 3-parameter with uniform rates among sites and a bootstrap of 100. Blue rhomboid indicates that the strain belongs to the 2000 collection Red circles to the 2010 collection. Green rectangle indicates 5 CRISPR arrays only detected in the 2010 strains.
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