



This is a repository copy of *A single natural nucleotide mutation alters bacterial pathogen host tropism*.

White Rose Research Online URL for this paper:  
<http://eprints.whiterose.ac.uk/108115/>

Version: Accepted Version

---

**Article:**

Viana, D., Comos, M., McAdam, P.R. et al. (8 more authors) (2015) A single natural nucleotide mutation alters bacterial pathogen host tropism. *Nature Genetics*, 47 (4). pp. 361-367. ISSN 1061-4036

<https://doi.org/10.1038/ng.3219>

---

**Reuse**

Unless indicated otherwise, fulltext items are protected by copyright with all rights reserved. The copyright exception in section 29 of the Copyright, Designs and Patents Act 1988 allows the making of a single copy solely for the purpose of non-commercial research or private study within the limits of fair dealing. The publisher or other rights-holder may allow further reproduction and re-use of this version - refer to the White Rose Research Online record for this item. Where records identify the publisher as the copyright holder, users can verify any specific terms of use on the publisher's website.

**Takedown**

If you consider content in White Rose Research Online to be in breach of UK law, please notify us by emailing [eprints@whiterose.ac.uk](mailto:eprints@whiterose.ac.uk) including the URL of the record and the reason for the withdrawal request.



[eprints@whiterose.ac.uk](mailto:eprints@whiterose.ac.uk)  
<https://eprints.whiterose.ac.uk/>

Published in final edited form as:

*Nat Genet.* 2015 April ; 47(4): 361–366. doi:10.1038/ng.3219.

## A single natural nucleotide mutation alters bacterial pathogen host-tropism

David Viana<sup>#1</sup>, María Comos<sup>#2</sup>, Paul R McAdam<sup>#3</sup>, Melissa J. Ward<sup>4</sup>, Laura Selva<sup>1</sup>, Caitriona M. Guinane<sup>3</sup>, Beatriz M. González-Muñoz<sup>5</sup>, Anne Tristan<sup>6</sup>, Simon J Foster<sup>5</sup>, J. Ross Fitzgerald<sup>3,10,\*</sup>, and José R. Penadés<sup>7,8,10,\*</sup>

<sup>1</sup>Universidad CEU Cardenal Herrera, 46113 Moncada, Valencia, Spain

<sup>2</sup>Centro de Investigación y Tecnología Animal, Instituto Valenciano de Investigaciones Agrarias(CITA-IVIA), Apdo. 187, 12.400 Segorbe, Castellón, Spain

<sup>3</sup>The Roslin Institute, University of Edinburgh, Easter Bush Campus, Edinburgh EH25 9RG, United Kingdom

<sup>4</sup>Centre for Immunity, Infection and Evolution, University of Edinburgh, Edinburgh, EH93JT

<sup>5</sup>Department of Molecular Biology and Biotechnology, University of Sheffield, Sheffield, S10 2TN, UK

<sup>6</sup>Centre National de Référence des Staphylocoques, Université Lyon, France

<sup>7</sup>Instituto de Biomedicina de Valencia (IBV-CSIC), 46010, Valencia, Spain

<sup>8</sup>Institute of Infection, Immunity and Inflammation, College of Medical, Veterinary and Life Sciences, University of Glasgow, Glasgow, G12 8TA, UK

# These authors contributed equally to this work.

### Abstract

The capacity of microbial pathogens to alter their host-tropism leading to epidemics in distinct host-species populations is a global public and veterinary health concern. In order to investigate the molecular basis of a bacterial host-switching event in a tractable host-species, we traced the evolutionary trajectory of the common rabbit clone of *Staphylococcus aureus*. We report that it evolved through a likely human-to-rabbit host jump over 40 years ago, and that only a single natural nucleotide mutation was required and sufficient to convert a human-specific *S. aureus*

---

Users may view, print, copy, and download text and data-mine the content in such documents, for the purposes of academic research, subject always to the full Conditions of use:[http://www.nature.com/authors/editorial\\_policies/license.html#terms](http://www.nature.com/authors/editorial_policies/license.html#terms)

\*Corresponding authors: José R. Penadés, Institute of Infection, Immunity and Inflammation, College of Medical, Veterinary and Life Sciences, University of Glasgow, JoseR.Penades@glasgow.ac.uk, Phone: +44 (0)141 330 8770; J. Ross Fitzgerald, The Roslin Institute and Edinburgh Infectious Diseases, University of Edinburgh, Ross.Fitzgerald@ed.ac.uk, Phone: +44 (0)131 6519235.

<sup>10</sup>These authors jointly supervised this work.

**Author Contributions.** J.R.F. and J.R.P. conceived and designed the study; D.V., M.C. and L.S. generated and characterized the different mutant strains; P.R.M, M.J.W. and C.M.G performed the genomic studies; B.M.G and S.J.F. measured D-Ala content; A.T. provided human strains; J.R.P, J.R.F and S.J.F supervised the research; J.R.F. and J.R.P. wrote the manuscript; J.R.F. and J.R.P. obtained funding.

**Accession codes.** The sequences reported in this paper have been deposited in the GenBank database (accession nos. ERS400826 to ERS400839). See Supplementary Table 1 for details.

**Competing Financial Interest.** The authors declare no competing financial interests.

strain into one which could infect rabbits. Related mutations were identified at the same locus in other rabbit strains of distinct clonal origin, consistent with convergent evolution. This first report of a single mutation that was sufficient to alter the host-tropism of a micro-organism during its evolution highlights the capacity of some pathogens to readily expand into novel host-species populations.

## Keywords

*Staphylococcus aureus*; host adaptation; evolution; virulence

---

## Introduction

The capacity for pathogens to switch host-species leading to epidemic spread in new host populations is a major veterinary and public health concern, but our understanding of the genetic basis of host-adaptation is very limited. Many animal- and plant-associated microorganisms co-evolve with a single host species leading to complex mutualistic, commensal or pathogenic relationships, and switches between host-species must overcome innate host-specific barriers to colonization and subsequent transmission. However, successful host jumps provide an opportunity to expand into novel ecological niches, and anthropogenic events such as domestication, industrialization of agriculture, and globalization have provided increased opportunities for the transmission of bacteria between humans and animals and their subsequent dissemination<sup>1</sup>. Recent studies have established that the host-shift and onward transmission of influenza virus requires several multi-gene mutations<sup>2-4</sup>, but the precise primary mechanisms have not been completely deciphered. The mechanisms underlying successful bacterial host-switching events are even less well understood but it is generally assumed that more complex genetic adaptations affecting multiple pathways would underlie adaptation to distantly related members of the animal kingdom.

*Staphylococcus aureus* is a major human pathogen that is also associated with economically important infections of livestock including dairy cows, sheep, poultry and rabbits<sup>5</sup>. Previously, it has been demonstrated that the major clones of *S. aureus* associated with ruminant and poultry infections evolved as the result of human to animal host jumps, leading to host-specialized endemic clones responsible for disease<sup>6-9</sup>. Several studies have demonstrated an association of specific mobile genetic elements (MGEs) with strains infecting particular host species suggestive of a role for MGE in host-specificity<sup>7,8,10-12</sup>. However, the molecular basis for *S. aureus* host-adaptation and the underlying genetic events involved is unclear.

## Results

### ***S. aureus* clonal complex CC121 has a multi-host tropism**

A common cause of human skin and soft tissue infections such as staphylococcal scalded skin syndrome, and life-threatening infections such as necrotizing pneumonia is the multi-locus sequence typing (MLST)-defined clone ST121<sup>13</sup>. Recently, the industrialization of

rabbit farming in the developed world has coincided with the emergence of a highly-virulent epidemic clone of *S. aureus*, also ST121, which is associated with skin abscess and mastitis infections of rabbits in commercial rabbitries<sup>14</sup>. However, the basis for the multiple host species association is unknown. We hypothesized that the multiple host association of ST121 strains could be explained either by the existence of host-specific subtypes of ST121, or an innate capacity of ST121 strains to infect both human and rabbit hosts. In order to address this question, we carried out experimental intradermal infections of rabbits with representative ST121 isolates of rabbit (n=3) and human (n=3) clinical origin. The rabbit ST121 strains caused infections resulting in skin abscesses which are characteristic of clinical symptoms seen in natural rabbit skin infections at doses as low as 300 cfu (Fig. 1a, b and c). In remarkable contrast, human ST121 strains failed to cause clinical infections with overt symptoms even at an inoculum of 10<sup>5</sup> cfu (Fig. 1a and b). These data indicate that in contrast to human ST121 isolates, rabbit ST121 strains have the capacity to cause disease in experimental infection of rabbits at very low inocula.

### **A human to rabbit *S. aureus* host jump occurred over 40 years ago**

Given the existence of a subset of ST121 strains with the capacity to cause clinical disease in rabbits, we investigated their evolutionary history. Accordingly, we compared the whole genome sequences from a total of 23 different ST121 strains which broadly represented the breadth of spatial and temporal diversity of the ST121 lineage including 14 newly-sequenced *S. aureus* ST121 isolates obtained from humans (n=8) and rabbits (n=6) in 8 countries (from 3 continents) over a 50 year period (Supplementary Table 1), in addition to 11 diverse publicly available genome sequences for human ST121 strains<sup>15</sup>. Of note, one of the rabbit strains (DL190) was sequenced to completion resulting in a whole genome sequence which could be used as a reference for comparison to all other ST121 sequences. We then applied a Bayesian coalescent method in the program Bayesian Evolutionary Analysis Sampling Trees (BEAST) employing a relaxed molecular clock model to reconstruct the phylogeny of the ST121 human and rabbit strains and estimate the rate of evolution of the ST121 clone<sup>16</sup>. The resulting phylogenetic tree indicates a high level of diversity of human strains represented by long branches in the ST121 phylogenetic tree in contrast to a single clade comprised of all rabbit strains which were isolated 20 years apart in 4 different countries (Fig. 2 and Supplementary Table 1). Using the calculated rate of molecular evolution for the ST121 sequences examined ( $1.83 \times 10^{-6}$  substitutions per site per year (95% HPDs  $9.84 \times 10^{-7}$  -  $2.26 \times 10^{-6}$ ), we estimated the date of the most recent common ancestor (MRCA) of the rabbit clade to be 1976 (1942-1990), and the MRCA of the rabbit clade with the most closely-related human strain F as 1909 (1804-1958). Taking into account the topology of the ST121 tree and the previously inferred human ancestry of the *S. aureus* species<sup>6</sup>, the most likely explanation for the existence of the ST121 rabbit clade is a single human-to-rabbit host jump which occurred more than 38 years ago leading to the emergence of a new clone associated with epidemics of rabbit farms (Fig. 2).

### **Diversification of the rabbit *S. aureus* ST121 core genome**

Considering previous studies which have suggested an important role for MGEs in adaptation to human, ruminant and avian hosts<sup>7,8,10-12</sup>, we hypothesised that the ST121 clone had adapted to infect rabbits using a similar strategy. Comparative analysis of the

accessory genome of ST121 strains revealed that the majority of human strains contained MGEs which encode potent toxins involved in human disease pathogenesis such as Pantone-Valentine leukocidin (PVL) and exfoliative toxins (ET), and all but one contained a  $\beta$ -converting phage encoding the human-specific immune evasion cluster (IEC)<sup>17</sup>. None of the rabbit strains contained PVL- or ET-encoding MGEs indicating their dispensability for *S. aureus* infections of rabbits (Fig. 2). Unexpectedly, the rabbit strains did not contain any MGE which are unique to rabbit *S. aureus* (Fig. 2) implying that the host tropism of the rabbit ST121 clade is not mediated by a niche-specific accessory gene pool as previously reported for *S. aureus* clones associated with poultry and ruminants<sup>5</sup>. Accordingly, we concluded that the capacity of ST121 to infect rabbits must have evolved through mutational events in the core genome of an ST121 progenitor of the rabbit clade. Core genome analysis revealed mutations common to all rabbit strains and absent from human strains including a total of 9 predicted pseudogenes resulting from non-sense mutations leading to premature stop codons or indels leading to frameshifts (Supplementary Table 2), and 254 non-synonymous (NS) mutations in a variety of gene loci (Supplementary Table 3). Of the genes associated with loss-of-function mutations, 4 had previously been implicated in *S. aureus* virulence including genes that would encode urease, lipase, EssB, and the global gene regulator *rot* (repressor of toxins)<sup>18-20</sup>. In addition, a single pseudogene caused by the insertion of a prophage into the  $\beta$ -toxin gene (*hlyB*) was identified in all human ST121 strains but was intact in the rabbit strains examined. Further, a SNP in the *dltB* gene encoding the D-alanine lipoteichoic acid and wall teichoic acid esterification protein resulted in the conversion of the stop codon into a tyrosine and extended the length of the predicted protein by a single amino acid residue. In addition, there were 2 further NS SNPs in the *DltB* coding sequence (Supplementary Fig. 1). Finally, an additional single 717 bp gene (*spE*) encoding a predicted serine protease was identified in each of the rabbit strains in the vSa $\beta$  genomic island (Supplementary Table 2).

### A single non-synonymous mutation is required for *S. aureus* rabbit infectivity

In order to distinguish host-adaptive mutational events among neutral or mildly deleterious mutations associated with genetic drift, mutations which were associated with encoded proteins of different size in all rabbit strains (ie resulting from frameshift, nonsense or length extension mutations) were restored in representative rabbit strains (strains I and J) and introduced into the human ST121 strain F by allele replacement. In addition, the single gene *spE* found in the rabbit strains was deleted in strains I and J. The infectivity and severity of experimental infection by wild type and derived mutant strains were then compared in a rabbit skin abscess infection model. For mutations in 10 of the 12 genes examined, reversion to the human ancestral gene state in the rabbit strains and their introduction to the human ST121 strain F had no effect on infectivity or severity (Supplementary Table 2). However, reversion of the 3 identified *dltB* SNPs (*dltB<sup>h</sup>*) in the rabbit strains J and I resulted in complete loss of infectivity, even at an infectious dose of 10<sup>5</sup> cfu (equivalent to a wild type human strain; Fig. 3a). In addition, reversion of the nonsense mutation in the global virulence regulator *rot* (*rot+*) significantly reduced the severity and infectivity of the rabbit strains (Fig. 3a).

In order to examine the role of specific *dltB* mutations in rabbit infectivity, each of the 3 *dltB* mutations, in addition to the *rot* nonsense mutation, were individually introduced to the human ST121 strain F. Remarkably, introduction of the most 5' NS SNP of the *dltB* locus in the rabbit strains (T113K) to human ST121 strain F, conferred the capacity to infect the rabbit host at an inoculum of 300 cfu (Fig. 3b), causing lesions that were highly similar to that caused by the wt rabbit strain J (Fig. 3c and d). To confirm that the infective phenotype of the ST121 strain F DltB T113K was not due to unrelated mutations which had occurred elsewhere in the genome during the synthesis of the mutant, we restored the *dltB* gene to its original wt state in strain F and observed the expected loss of infectivity. In contrast, neither of the other 2 SNPs in the *dltB* gene (Y250H and 405Y) nor the *rot* mutation (*rot*-) increased the infectivity of the human clone F when introduced individually. Of note, the introduction of DltB T113K resulted in lesions in 40% of inoculated animals but bacteria were recovered from only about 20%. However, the introduction of all 3 *dltB* mutations resulted in infected lesions with 100% recovery, with an infectivity which is equivalent to the rabbit strain with a functional *rot* gene (human ancestral state) (Fig 3 b). These data suggest that additional *dltB* mutations promote persistence of *S. aureus* ST121 in infected lesions. Importantly, a combination of the *rot* nonsense mutation with the 3 *dltB* SNPs in the human F strain resulted in a strain whose infectivity was indistinguishable from the wt rabbit strains resulting in lesions with highly similar gross and microscopic histopathology (Fig. 3b). Finally, we also verified that the rabbit strain J with a deleted *rot* gene demonstrated no further attenuation for infectivity, suggesting that loss of *rot* gene function rather than allelic diversification of the *rot* gene is responsible for the attenuation. Taken together, the data indicate that a single natural nucleotide mutation of *dltB* was sufficient to confer the capacity of *S. aureus* ST121 to cause intradermal infections of rabbits, and that 2 additional NS mutations of *dltB* and loss of function of *rot* elevated the infectivity and bacterial recovery to that of wild type *S. aureus* rabbit strains.

### DltB mutation-related D-Ala levels do not correlate with rabbit infectivity

DltB is an integral membrane protein encoded by the *dltABCD* operon which is likely responsible for the translocation and incorporation of D-alanine into teichoic acids and lipoteichoic acids of *S. aureus*<sup>21</sup>. While our understanding of the function of teichoic acids is incomplete, the addition of D-alanine (D-Ala) residues confers a positive charge which is essential for resistance to cationic peptides and for virulence<sup>21,22</sup>. In view of this, we speculated that the activity of the mutated form of DltB may be enhanced leading to increased D-Ala substitution. Importantly, we confirmed that the variant DltB protein from rabbit strain J is fully functional, as a *dltB* deletion mutant was incapable of infecting the rabbit at a dose of 10<sup>5</sup> cfu. Although D-alanylation of wall teichoic acids as well as resistance to the cationic peptide-mediated killing were increased in the rabbit strain compared to the human clone, this phenotype was not dependent on the *dltB* mutations (Supplementary Fig. 2a and 2b). In addition, there was no difference in D-Ala levels associated with lipoteichoic acids from strains with the different *dltB* alleles (Supplementary Fig. 2c). Considering the finding that D-Ala content does not have a central impact on rabbit infectivity, we analysed if the DltB mutations affected the bacterial cell wall composition. As shown in Supplementary Figure 3a and 3b, there were no differences in the structure or amino acid composition of the peptidoglycan obtained from the different strains. Next,

considering that the DltB mutations present in the rabbit clone are predicted to have an extracellular or proximal extracellular location (Supplementary Fig. 4), we analysed if they could be involved in interacting with the innate immune system of the host or with other components of the bacterial surface. This was addressed by measuring survival in a whole blood killing assay, and by analysing the protein profile of the proteins bound to the bacterial surface. As shown in Supplementary Figure 5 and 6, although there were phenotypic differences between the human and the rabbit clones, they were not attributable to the DltB mutations. Of note is the data from the killing assays showing an increased resistance of the F *dltB*<sup>r</sup> clone to killing compared to the wild-type F clone (Supplementary Fig. 5). However, since the mutated rabbit clone J *dltB*<sup>h</sup> which is uninfecive for rabbits did not demonstrate reduced resistance to killing compared to that observed with the J wild-type, and the survival of the J *dltB*<sup>h</sup> clone in blood was higher to that observed with the F *dltB*<sup>r</sup> clone, it was concluded that the ability to infect rabbits does not likely depend on the resistance observed in the killing assay. In conclusion, these data demonstrate that although we detected phenotypic differences between the human and the rabbit clones, they are not dependent on the DltB mutations. This implies that the effect of the mutated DltB on rabbit infectivity is due to alternative functions of DltB during infection. In support of this, it has been recently proposed that DltB is a member of a new superfamily of proteins, named MBOAT (for membrane-bound O-acyl transferases), that transfer organic acids, typically fatty acids, onto hydroxyl groups, performing a role in signalling<sup>23</sup>. However, to date, O-acyl transferase enzymatic activity has not been demonstrated experimentally for DltB, and the mechanism underlying the role of the *dltB* mutations in adaptation to rabbits is unclear.

### Convergent evolution of *S. aureus* at the *dltB* locus

Finally, we hypothesized that if a single nucleotide mutation of *dltB* was essential to change the host tropism of ST121, that similar mutations will have occurred in other *S. aureus* strains infecting rabbits and perhaps other bacterial species. Phylogenetic analysis of rabbit *S. aureus* strains based on concatenated MLST sequences indicates that the capacity to infect rabbits has occurred on numerous occasions, presumably through human-to-rabbit host jump events (Fig. 4). In order to investigate the possibility that the adaptive mutation associated with the *dltB* locus in ST121 rabbit strains is a widespread *S. aureus* rabbit host-adaptive mechanism, we sequenced the *dltB* gene of representative rabbit isolates from *S. aureus* ST1, ST8, ST9, ST45, ST96, ST133 and ST398, lineages which each include isolates of both human and rabbit clinical origin (table 1 and Fig. 4), and compared them to *dltB* sequences from diverse human and animal isolates. DltB was highly conserved among the great majority of human, ruminant and poultry strains of *S. aureus* with only 8 of 445 strains examined (1.8%) containing one or more NS mutations (Supplementary Fig. 1). However, 39 of 39 (100%) rabbit strains examined from diverse clonal lineages contained one or more NS SNPs in *dltB*, suggestive of convergent evolution (Table 1 and Fig. 4). Of note, the majority of the mutations are also predicted to be extracellularly exposed or proximal to the outer surface of the membrane (Supplementary Fig. 4). In order to further examine the convergent host-adaptive evolution of rabbit *S. aureus* clones, we tested the role in virulence of the distinct *dltB* allele associated with the *S. aureus* ST96 rabbit clone (differing at 2 amino acid residues, Table 1). Introduction of the ST96 rabbit *dltB* allele to rabbit ST121 conferred a similar level of infectivity to the ST121 rabbit *dltB* allele (10 out of 15 animals

infected with the ST121 allele *vs* 7 out of 15 animals infected with the ST96 allele;  $p=0.46$ , Yates' chi-squared test). Taken together, these data support the conclusion that distinct *dltB* mutations occurring in different *S. aureus* strain backgrounds underpinned independent host-jump events from humans to rabbits. In contrast to the allelic diversity of *dltB* identified among rabbit strains, all rabbit strains except ST121 contained an intact *rot* gene, and the majority contained the  $\beta$ -converting phage, resulting in a non-functional  $\beta$ -toxin, indicating that mutations affecting those determinants in ST121 are not essential for rabbit host-adaptation. Taken together, the data are consistent with the critical role for mutations of *DltB* in *S. aureus* rabbit host adaptation.

Finally, we identified strains of other bacterial species which had non-synonymous *dltB* mutations and predicted extended *DltB* proteins. These included a soil-adapted subtype of the plant bacterium *Bacillus amyloliquefaciens* which has several predicted non-synonymous mutations and an additional histidine residue at the *DltB* C-terminus, that were absent in the plant root-adapted strain (Supplementary Fig. 7a). In addition, analysis of publicly available *Streptococcus pneumoniae* genome sequences from a study examining resistance to vaccine and therapeutic pressures in humans, revealed additional predicted residues of *DltB* proteins in several unrelated strains (Supplementary Fig. 7b). Taken together, these data suggest a potential general role for allelic variants of *DltB* proteins in adaptation to specific environmental or host-specific niches.

## Discussion

The molecular basis of the host-adaptation of viral pathogens such as influenza and HIV has been intensively investigated. Several key mutations such as influenza PB2-627 and HIV Gag-30 have been identified as sites of critical adaptive mutations<sup>24,25</sup>. However, the adaptation to mammalian hosts by both influenza and HIV appears to involve multi-gene mutations<sup>26,27</sup>. For bacterial host adaptation, horizontal acquisition of a single gene regulator in the bacterial squid symbiont *Vibrio fischeri* was demonstrated to be an essential step in adaptation to its host<sup>28</sup>. In addition, previous seminal work demonstrated that in a mouse model of *Listeria monocytogenes* infection, a single amino acid change in the mouse E-cadherin receptor increased *L. monocytogenes* cell internalisation *in vitro*<sup>29</sup>, and conversely, 2 amino acid changes in the listerial invasion protein InlA increased *in vivo* infectivity by enhancing bacterial affinity to the murine E-cadherin<sup>30</sup>. However, very high inoculum doses were still required for infectivity, suggesting that for natural *L. monocytogenes* infections to occur additional adaptations would be required to facilitate a host shift.

To date, the fundamental biological question of how animal-bacteria partnerships are established has been difficult to dissect via established animal models of infection. In the current study, the fact that the bacterial host jump being examined involved rabbits, an experimentally-tractable host-species, facilitated for the first time, a dissection of the evolutionary genetic trajectory of a natural host-species switch by a bacterial pathogen. Using this approach, we traced the evolutionary genetic events which led to the emergence of a major animal clone of *S. aureus* responsible for disease epidemics of farmed rabbits on a global scale. In contrast to all characterized human and animal clones of *S. aureus*,



adaptation to the rabbit host did not involve acquisition of MGE from a host-specific accessory gene-pool. Remarkably, we report that a single nucleotide mutation, which occurred naturally, was sufficient to allow a radical change in bacterial host-tropism. The mutation, affecting the *dllB* gene was sufficient and required to convert a human *S. aureus* strain, which was incapable of causing infections in rabbits, to one that had the capacity to cause epidemics in farmed rabbit populations. An additional 2 mutations of the same locus enhanced infectivity and bacterial proliferation within rabbit skin lesions. In spite of apparently numerous host jump events into rabbits, ST121 is by far the dominant clone in the countries sampled. As such, we suggest that in addition to the initial adaptations required for survival in the new host species, additional mutations likely contribute to the transmissibility and successful spread of the infectious clone.

In summary, our results reporting a single natural nucleotide mutation associated with a bacterial host switch event represent a paradigm shift in understanding of the minimal adaptations required for a bacterium to overcome species-barriers and establish in new host populations. The discovery has important public and veterinary health implications which will require a re-examination of the future threat posed by bacterial pathogen host switching events.

## METHODS

### Bacterial strains and growth conditions

Bacterial strains used are listed in Tables S1 and S4. *S. aureus* strains were grown at 37°C on TSA agar medium, supplemented with antibiotics as appropriate. Broth cultures were grown at 37°C in TSB broth with shaking (240 r.p.m.). LB agar and LB medium were used for *E. coli* strains, supplemented with antibiotics as appropriate. Procedures for transduction and transformation in *S. aureus* were performed as described previously<sup>31,32</sup>.

### DNA methods

General DNA manipulations were performed by standard procedures. To introduce specific mutations into *S. aureus* strains, we used plasmid pMAD<sup>33</sup> for allelic exchange as previously described<sup>31</sup>. Plasmid constructs (listed in Table S5) were prepared by cloning PCR products obtained with oligonucleotide primers as listed in Table S6, and all clones were sequenced by the Institute Core Sequencing Lab.

### Whole genome sequencing and alignment

Isolates were sequenced using a combination of single and paired end technologies on the Roche 454 platform and assembled using the Roche GS De Novo Assembler v2.6, or alternatively using the Illumina HiSeq with reads assembled using Velvet v1.2.08. Contigs were aligned using the progressiveMauve algorithm, filtered for locally colinear blocks over 1000 nucleotides in length, and gap positions were removed<sup>34</sup>. The alignment was assessed for recombination using BRATNextGen<sup>35</sup>, with predicted recombinant regions excluded from further phylogenetic analyses. Genes specific to rabbit ST121 isolates were identified by comparing the sequenced rabbit isolate genome sequences to the human ST121 sequences using the `print_novel_contigs` function of `cortex_var` v1.0.5.20<sup>36</sup>.

### Bayesian evolutionary analysis

Based on the core genome Mauve alignment (6208 variable sites), the phylogeny of ST121 was reconstructed using BEAST v1.6.1<sup>16</sup> under the GTR model of nucleotide substitution with a gamma correction for rate heterogeneity and a skyline coalescent tree prior. Simultaneously we used an uncorrelated lognormal distribution to model the rate of evolution and constrained the tips of the phylogeny to their dates of isolation to calibrate the rate<sup>37</sup>. Posterior probabilities were estimated from 3 independent Markov chain Monte Carlo (MCMC) samples from 3 independent analyses each run for  $1 \times 10^8$  iterations, with sampling at every 1000 generations and 10% discarded as burnin.

### Identification of pseudogenes and non-synonymous mutations

Single end 454 sequence reads were aligned to the ED133 genome (accession number NC\_017337.1), and an annotated draft version of the DL190 genome using the BWA long read aligner<sup>38</sup>. Point mutation and insertion/deletion variants were called at sites with average mapping and base quality scores greater than 30. Mutations predicted to result in pseudogenes were further inspected manually using the Integrative Genomics Viewer<sup>39</sup>. Variation in gene content between strains was examined using the GeneFamily method of the Pangenome Analysis Pipeline with default parameters<sup>40</sup>. Presence or absence of virulence factors was confirmed through manual inspection of the output of BLAST searches.

### Rabbit skin infection model

Two-month old albino hybrid rabbits of either sex were used for the skin infection model. Rabbits were sedated with a combination of ketamine and xylazine and were shaved on their back and inoculated by intradermal injection (usually with 300 cfu of *S. aureus* in 100  $\mu$ l of PBS). Each rabbit was inoculated in duplicate with 2 strains (generally wild type and mutant), and each strain was tested in groups of at least 20 rabbits. After 7 d, rabbits were euthanized by intravenous injection of barbiturate (Dolethal, Vétquinol SA). In each experiment, an additional group of animals was inoculated with vehicle (PBS) and served as a negative control. In addition, to exclude the possibility of contamination, bacteria recovered at the end of the experimental period were analysed by sequencing of strain-specific gene alleles. The experimental protocol was approved by the ethical committee of the Universidad CEU Cardenal Herrera and by the Conselleria d' Agricultura, Pesca i Alimentació, Generalitat Valenciana (permit number 2011/010).

### Macroscopical and histological examination of lesion tissues

Rabbits were examined daily and development of infection was followed visually and by palpation of lesion tissue. Length and width values were measured to calculate the area of lesions. Addition of lesion size, skins were also examined to evaluate presence or absence of epidermal necrosis. After post-mortem examination, skin abscesses were routinely processed for histological examination and stained with haematoxylin and eosin and by Gram's method. Previously fixation, swab samples were taken from lesion and cultivated on blood-agar (BioMérieux, Marcy l' Etoile, France) to confirmed presence of *S. aureus*.

## Cell wall and D-alanine analyses

Cell walls were isolated, wall teichoic acids and lipoteichoic acids were purified and D-alanine content measured as previously described<sup>21,41</sup>. The peptidoglycan composition and structure in the different strains was analysed by Cocolabs (Tuebingen, Germany), as described before<sup>42</sup>.

## Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

## Acknowledgements

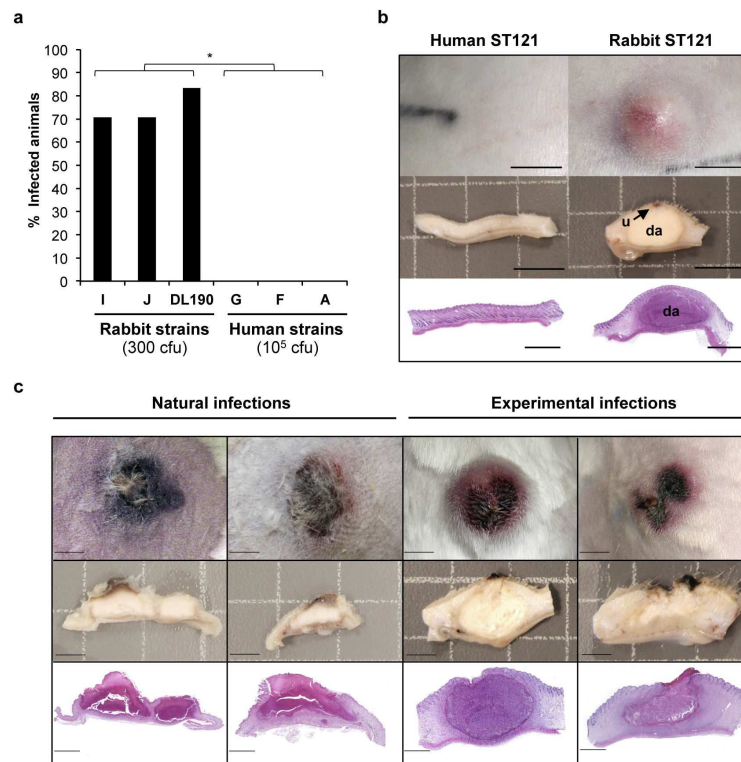
We thank Jerome Etienne for helpful advice, Olaf Schneewind, Mark Woolhouse and Íñigo Lasa for comments on the manuscript, Concha Cervera and Enrique Blas for their support with the *in vivo* experiments, and Robyn Cartwright for excellent technical assistance. This work was supported by grants BIO2011-30503-C02-01, Eranet-pathogenomics PIM2010EPA-00606 and Consolider-Ingenio CSD2009-00006 from Ministerio de Ciencia e Innovación (Spain), and strategic grant funding from University of Glasgow to J.R.P; by a project grant (BB/I013873/1) and institute strategic grant funding from the Biotechnology and Biological Sciences Research Council (United Kingdom) to J.R.F., in addition to a doctoral training grant from the Medical Research Council (UK) to J.R.F; and by grants AGL2011-30170-CO2-02 (Ministerio de Ciencia e Innovación, Spain) and GV2013-077 (Conselleria d' Educació, Cultura i Esport, GV) to D.V. We are grateful to Edinburgh Genomics (Roslin Institute) for sequencing services.

## References

1. Marano N, Arguin PM, Pappaioanou M. Impact of globalization and animal trade on infectious disease ecology. *Emerging Infect. Dis.* 2007; 13:1807–1809. [PubMed: 18258027]
2. Imai M, et al. Experimental adaptation of an influenza H5 HA confers respiratory droplet transmission to a reassortant H5 HA/H1N1 virus in ferrets. *Nature.* 2012; 486:420–428. [PubMed: 22722205]
3. Russell CA, et al. The potential for respiratory droplet-transmissible A/H5N1 influenza virus to evolve in a mammalian host. *Science.* 2012; 336:1541–1547. [PubMed: 22723414]
4. Herfst S, et al. Airborne transmission of influenza A/H5N1 virus between ferrets. *Science.* 2012; 336:1534–1541. [PubMed: 22723413]
5. Fitzgerald JR. Livestock-associated *Staphylococcus aureus*: origin, evolution and public health threat. *Trends Microbiol.* 2012; 20:192–198. [PubMed: 22386364]
6. Weinert LA, et al. Molecular dating of human-to-bovid host jumps by *Staphylococcus aureus* reveals an association with the spread of domestication. *Biol. Lett.* 2012; 8:829–832. [PubMed: 22628096]
7. Guinane CM, et al. Evolutionary genomics of *Staphylococcus aureus* reveals insights into the origin and molecular basis of ruminant host adaptation. *Genome Biol Evol.* 2010; 2:454–466. [PubMed: 20624747]
8. Lowder BV, et al. Recent human-to-poultry host jump, adaptation, and pandemic spread of *Staphylococcus aureus*. *Proc. Natl. Acad. Sci. U.S.A.* 2009; 106:19545–19550. [PubMed: 19884497]
9. Price LB, et al. *Staphylococcus aureus* CC398: host adaptation and emergence of methicillin resistance in livestock. *MBio.* 2012; 3
10. Viana D, et al. Adaptation of *Staphylococcus aureus* to ruminant and equine hosts involves SaPI-carried variants of von Willebrand factor-binding protein. *Mol. Microbiol.* 2010; 77:1583–1594. [PubMed: 20860091]
11. van Wamel WJB, Rooijackers SHM, Ruyken M, Van Kessel KPM, Van Strijp JAG. The innate immune modulators staphylococcal complement inhibitor and chemotaxis inhibitory protein of *Staphylococcus aureus* are located on beta-hemolysin-converting bacteriophages. *J. Bacteriol.* 2006; 188:1310–1315. [PubMed: 16452413]

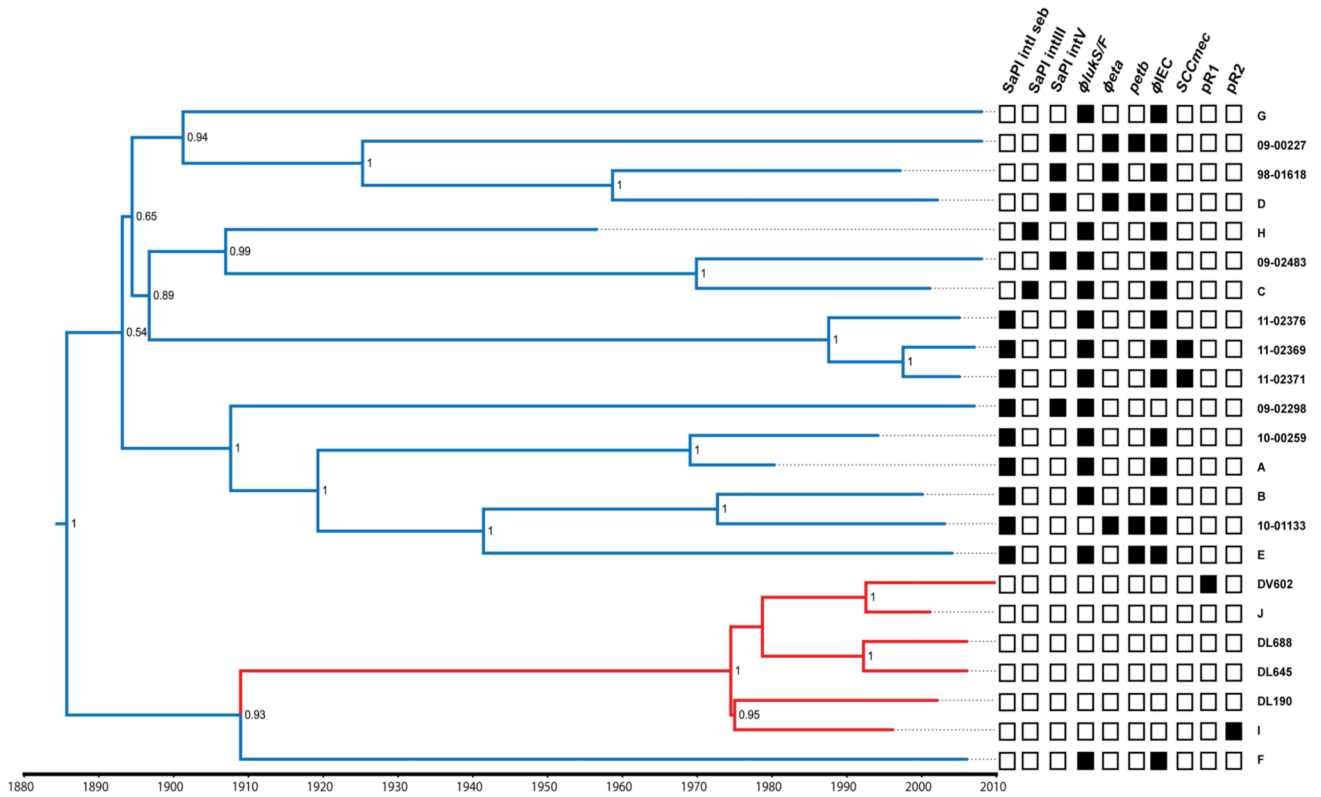
12. Ubeda C, et al. Sip, an integrase protein with excision, circularization and integration activities, defines a new family of mobile *Staphylococcus aureus* pathogenicity islands. *Mol. Microbiol.* 2003; 49:193–210. [PubMed: 12823821]
13. Rasigade JP, et al. Global distribution and evolution of Pantone-Valentine leukocidin-positive methicillin-susceptible *Staphylococcus aureus*, 1981-2007. *J. Infect. Dis.* 2010; 201:1589–1597. [PubMed: 20367458]
14. Vancraeynest D, et al. International dissemination of a high virulence rabbit *Staphylococcus aureus* clone. *J. Vet. Med. B Infect. Dis. Vet. Public Health.* 2006; 53:418–422. [PubMed: 17062118]
15. Kurt K, et al. Subpopulations of *Staphylococcus aureus* clonal complex 121 are associated with distinct clinical entities. 2013; 8:e58155.
16. Drummond AJ, Rambaut A. BEAST: Bayesian evolutionary analysis by sampling trees. *BMC Evol. Biol.* 2007; 7:214. [PubMed: 17996036]
17. Spaan AN, et al. The staphylococcal toxin Pantone-Valentine Leukocidin targets human C5a receptors. *Cell Host Microbe.* 2013; 13:584–594. [PubMed: 23684309]
18. Burts ML, Williams WA, DeBord K, Missiakas DM. EsxA and EsxB are secreted by an ESAT-6-like system that is required for the pathogenesis of *Staphylococcus aureus* infections. *Proc. Natl. Acad. Sci. U.S.A.* 2005; 102:1169–1174. [PubMed: 15657139]
19. Hu C, Xiong N, Zhang Y, Rayner S, Chen S. Functional characterization of lipase in the pathogenesis of *Staphylococcus aureus*. *Biochem. Biophys. Res. Commun.* 2012; 419:617–620. [PubMed: 22369949]
20. Saïd-Salim B, et al. Global regulation of *Staphylococcus aureus* genes by Rot. *J. Bacteriol.* 2003; 185:610–619. [PubMed: 12511508]
21. Peschel A, et al. Inactivation of the *dlt* operon in *Staphylococcus aureus* confers sensitivity to defensins, protegrins, and other antimicrobial peptides. *J. Biol. Chem.* 1999; 274:8405–8410. [PubMed: 10085071]
22. Collins LV, et al. *Staphylococcus aureus* strains lacking D-alanine modifications of teichoic acids are highly susceptible to human neutrophil killing and are virulence attenuated in mice. *J. Infect. Dis.* 2002; 186:214–219. [PubMed: 12134257]
23. Hofmann K. A superfamily of membrane-bound O-acyltransferases with implications for wnt signaling. *Trends Biochem. Sci.* 2000; 25:111–112. [PubMed: 10694878]
24. Subbarao EK, London W, Murphy BR. A single amino acid in the PB2 gene of influenza A virus is a determinant of host range. *J. Virol.* 1993; 67:1761–1764. [PubMed: 8445709]
25. Wain LV, et al. Adaptation of HIV-1 to its human host. *Mol. Biol. Evol.* 2007; 24:1853–1860. [PubMed: 17545188]
26. Naffakh N, Tomoiu A, Rameix-Welti M-A, van der Werf S. Host restriction of avian influenza viruses at the level of the ribonucleoproteins. *Annu. Rev. Microbiol.* 2008; 62:403–424. [PubMed: 18785841]
27. Etienne L, Hahn BH, Sharp PM, Matsen FA, Emerman M. Gene loss and adaptation to hominids underlie the ancient origin of HIV-1. *Cell Host Microbe.* 2013; 14:85–92. [PubMed: 23870316]
28. Mandel MJ, Wollenberg MS, Stabb EV, Visick KL, Ruby EG. A single regulatory gene is sufficient to alter bacterial host range. *Nature.* 2009; 458:215–218. [PubMed: 19182778]
29. Lecuit M, et al. A single amino acid in E-cadherin responsible for host specificity towards the human pathogen *Listeria monocytogenes*. *EMBO J.* 1999; 18:3956–3963. [PubMed: 10406800]
30. Wollert T, et al. Extending the host range of *Listeria monocytogenes* by rational protein design. *Cell.* 2007; 129:891–902. [PubMed: 17540170]
31. Tormo-Más MÁ, et al. Moonlighting bacteriophage proteins derepress staphylococcal pathogenicity islands. *Nature.* 2010; 465:779–782. [PubMed: 20473284]
32. Tormo-Más MÁ, et al. Phage dUTPases control transfer of virulence genes by a proto-oncogenic G protein-like mechanism. *Mol. Cell.* 2013; 49:947–958. [PubMed: 23333307]
33. Arnaud M, Chastanet A, Débarbouillé M. New vector for efficient allelic replacement in naturally nontransformable, low-GC-content, gram-positive bacteria. *Appl. Environ. Microbiol.* 2004; 70:6887–6891. [PubMed: 15528558]

34. Darling AE, Mau B, Perna N. T. progressiveMauve: multiple genome alignment with gene gain, loss and rearrangement. *PLoS ONE*. 2010; 5:e11147. [PubMed: 20593022]
35. Marttinen P, et al. Detection of recombination events in bacterial genomes from large population samples. *Nucleic Acids Res*. 2012; 40:e6–e6. [PubMed: 22064866]
36. Iqbal Z, Turner I, McVean G. High-throughput microbial population genomics using the Cortex variation assembler. *Bioinformatics*. 2013; 29:275–276. [PubMed: 23172865]
37. Drummond AJ, Ho SYW, Phillips MJ, Rambaut A. Relaxed phylogenetics and dating with confidence. *PLoS Biol*. 2006; 4:e88. [PubMed: 16683862]
38. Li H, Durbin R. Fast and accurate long-read alignment with Burrows-Wheeler transform. *Bioinformatics*. 2010; 26:589–595. [PubMed: 20080505]
39. Robinson JT, et al. Integrative genomics viewer. *Nat. Biotechnol*. 2011; 29:24–26. [PubMed: 21221095]
40. Zhao Y, et al. PGAP: pan-genomes analysis pipeline. *Bioinformatics*. 2012; 28:416–418. [PubMed: 22130594]
41. Gründling A, Schneewind O. Synthesis of glycerol phosphate lipoteichoic acid in *Staphylococcus aureus*. *Proc. Natl. Acad. Sci. U.S.A.* 2007; 104:8478–8483. [PubMed: 17483484]
42. Bertsche U, et al. Increased cell wall teichoic acid production and D-alanylation are common phenotypes among daptomycin-resistant methicillin-resistant *Staphylococcus aureus* (MRSA) clinical isolates. *PLoS ONE*. 2013; 8:e67398. [PubMed: 23785522]



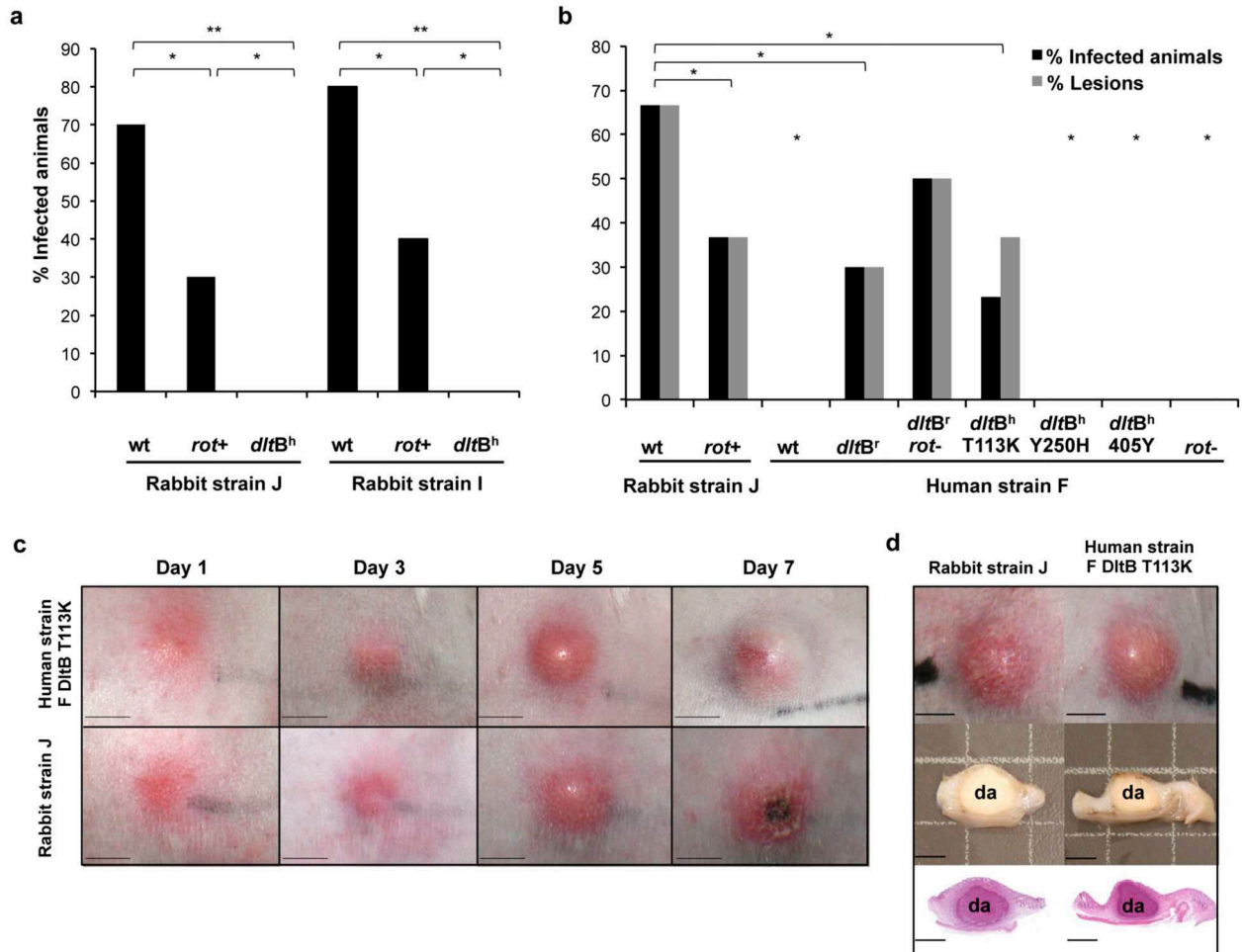
### Figure 1. Rabbit but not human ST121 strains can infect rabbits

(a) Percentage of rabbits infected with the rabbit or human ST121 clones (day 7 after inoculation). Rabbits ( $n = 24$  per strain) were inoculated intradermally with either *S. aureus* ST121 rabbit strains I, J or DL190 (300 cfu) or human strains G, F or A ( $10^5$  cfu), as described in Methods. Yates' chi-squared test was used to compute P values for between-group comparisons; differences that are statistically significant are indicated by an asterisk ( $P < 0.0001$ ); all other comparisons were non significant. (b) Representative rabbit skin lesions (day 7 post-inoculation). Images of gross and microscopic histopathology are presented for representative animals as follows: panel 1, gross skin pathology. Bar = 1cm; panel 2, transversal section of skin lesions. Lesions resulting from infection with rabbit strains were characterized by dermal abscesses (da) up to 2 cm in diameter, with epidermal ulcers (u) in the most severe cases. Bar = 1cm; panel 3, H&E-stained sections of the skin inoculated with human or rabbit ST121 clones, respectively. Bar = 0.5 cm. Microscopically, areas of purulent material surrounded by fibrosis infiltrated by inflammatory cells were observed. (c) Experimental skin lesions (day 10 post-inoculation) were indistinguishable from natural lesions. Panel 1, gross skin pathology; panel 2, transversal section of skin lesions. panel 3, H&E-stained sections of the skin. Bar = 0.5 cm.



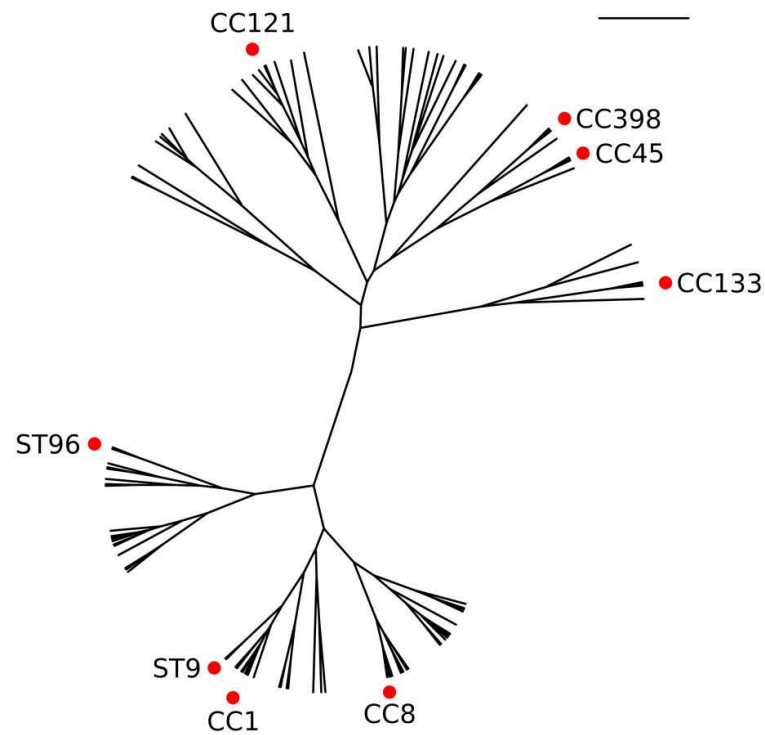
**Figure 2. Evolutionary history of the ST121 clonal complex suggests a human-to-rabbit host jump leading to the emergence of an epidemic rabbit-specific clone**

Bayesian phylogenetic reconstruction of the CC121 lineage based on core genome alignment with branches color-coded according to host species association (blue, human; red, rabbit). The presence or absence of mobile genetic elements (MGE) in the accessory genome is denoted by filled or empty squares, respectively. MGE identified include Staphylococcal Pathogenicity Islands (SaPI), phages ( $\phi$ ), or plasmids (p) containing genes: *lukF/S*, Panton-Valentine leukocidin; *eta*, exfoliative toxin A; *etb*, exfoliative toxin B; *seb*, staphylococcal enterotoxin B; IEC, immune evasion cluster; *SCCmec*, staphylococcal cassette chromosome *mec*, and plasmids encoding resistance to bleomycin, kanamycin, quaternary ammonium compounds and trimethoprim (pR1) or teicoplanin (pR2), respectively. Branch lengths are scaled according to the time-scale indicated on the x-axis.



**Figure 3. A single mutation is sufficient to confer rabbit infectivity to a human *S. aureus* strain**  
 Percentage of rabbits infected with the different rabbit (a) or human (b) ST121 mutants (day 7 after inoculation; n = 20 per strain in panel a, n = 30 per strain in panel b). Panel (b) also shows the percentage of animals that developed skin lesions by the inoculated bacteria. Yates' chi-squared test was used to compute P values for between-group comparisons; differences that are statistically significant are indicated by an asterisk (P < 0.05); all other comparisons were non significant. (c) The evolution of the skin lesions produced by the rabbit strain J and by the human clone carrying the DltB T113K mutation was indistinguishable. Rabbits were inoculated intradermally with 300 cfu of either the rabbit wt or the human mutant clones, as described in Methods. A representative animal is shown. The earliest gross changes were observed 24-48h post-infection (p.i.), consisting in a slight increase in size and erythema, but evolving to form skin abscesses up to 2 cm in diameter (7 day p.i.). Bar = 1,5 cm. (d) Transversal section of the skin lesions. Lesions produced by the rabbit and human mutant clones were characterized by dermal abscesses (da) up to 2 cm in diameter. Bottom: H&E-stained sections of the skins inoculated with the rabbit or human mutant ST121 clones. Bar = 0.5 cm. Microscopically, areas of purulent material surrounded by fibrosis infiltrated by inflammatory cells were observed.





**Figure 4. Rabbit *S. aureus* clones have evolved on numerous occasions and are associated with NS mutations of *dltB***

Bayesian phylogeny of *S. aureus* species reconstructed using nucleotide sequence of five non-recombinant MLST loci from 108 STs representing the breadth of species diversity. Clonal complexes/sequence types with known rabbit host association are indicated with red circles, and scale bar represents 5 nucleotides.

**Table 1**  
**Non-synonymous polymorphisms identified in the *dltB* locus from rabbit associated clonal complexes**

Clonal Complex	Sequence Type	Nucleotide change <sup>a</sup>	Non-synonymous polymorphisms	
1	1	680T>C	Ile227Thr	
		1202G>A	Gly401Asp	
	2780	1	5T>C	Ile2Thr
			1213T>C	*405Gln
			1213T>C	*405Gln
8	8	680T>G	Ile227Ser	
		1213T>C	*405Gln	
	407	363C>A	His121Gln	
		680T>C	Ile227Thr	
		1213T>C	*405Gln	
	879	363C>A	His121Gln	
		680T>C	Ile227Thr	
1213T>C		*405Gln		
9	9	1202G>A	Gly401Asp	
45	45	182T>G	Val61Gly	
		1037A>C	Tyr346Ser	
96	96	1205A>G	Lys402Arg	
		1213T>C	*405Gln	
121	121	338C>A	Thr113Lys	
		748T>C	Tyr250His	
		1215A>T	*405Tyr	
133	133	692A>G	Gln231Arg	
		739T>C	Tyr247His	
		1036T>C	Tyr346His	
		1202G>A	Gly401Asp	
398	398	680T>G	Ile227Ser	
		1213T>C	*405Gln	
	398	1037A>G	Tyr346Cys	

<sup>a</sup>In comparison with *dltB* from human-associated *S. aureus*.