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3 **Management of antibiotic resistance crisis: a**
4 **new horizon in climate change era**
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9 **ABSTRACT**
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The problem of antibiotic resistance develops when bacteria are able to grow in the presence of conventional antimicrobial drugs and today represents a serious public health issue. The environmental effects of global warming, by unknown genomic mechanisms of adaption, could dramatically increase this phenomenon and support a more rapid progression to “post-antibiotic era” in which common infections will be untreatable. Alternative approaches toward drug-resistant bacterial infections need to be explored to ensure effective therapies. Bacterial pathogens produce virulence factors, molecules that allow them to invade and to damage host cells. Methionine sulfoxide reductase enzymes are important, but poorly studied, virulence factors for many bacterial strains. A well knowledge in their mechanism of action and regulation could help in developing novel therapeutic strategies toward drug-resistant bacteria, in order to overcome the antibiotic resistance crisis.

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12 *Keywords: Antibiotic crisis; Bacterial infection, Global warming, Methionine sulfoxide reductase; Public health, Virulence*
13 *factor*
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16 **1. BACKGROUND**
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18 The first documented antibiotic resistance concerned the penicillin, that was discovered by Alexander Fleming in 1928,
19 and dates back to the 1940s, well before the use on a large scale that began during World War II. The proportions of
20 antibiotic resistance have grown exponentially over the past few years, leading to reduced therapeutic efficacy, and
21 increased mortality rates [1]. This emergency is recognized by the World Health Organization as one of the most
22 important public health threats affecting humans worldwide in this century. Current estimates suggest that by 2050 ten
23 million of premature deaths annually will be caused by resistant infections [2].

24 Antibiotic resistance is amplified by overuse or inappropriate prescription of antibiotics, the extensive use of them as
25 growth supplements in livestock, and the stall in development of new antibiotics by the pharmaceutical industry [1].
26 Climate change has been identified by the World Health Organization as a major factor in the spread of emerging
27 infectious disease worldwide. Climatic factors such as temperature, precipitation, and humidity modulate many biological
28 aspects concerning the transmission of pathogens [3]. More recently, a relationship between increased antibiotic
29 resistance of certain bacterial strains and global warming was observed [4]. The mechanism behind the phenomenon is
30 still unknown and, in view of this scenario, the impact of antibiotic resistance on global health could be dramatically
31 underestimated [5].

32 Bacterial adaptive response to antibiotics originates from the massive genetic plasticity of prokaryotic cells, based on
33 processes such as acquisition of genetic material through horizontal gene transfer, and alteration of gene expression.
34 These mechanisms of mutational adaption, an example of Darwinian principle of evolution, can confer to the pathogen
35 resistance to virtually all drugs currently available in clinical practice [6]. This aspect is so important that the term
36 “resistome” has been coined to define the set of genes that provide bacteria with an arsenal of weapons to resist
37 antibiotics. Furthermore, an open source database has been implemented (Comprehensive Antibiotic Resistance

38 Database, <http://arpcard.mcmaster.ca>), containing high quality reference data on the molecular basis of antimicrobial
39 resistance [7].

40 2. NEW PERSPECTIVES

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42 The knowledge of the biochemical and genetic basis of this phenomenon is fundamental to design novel therapeutic
43 strategies against antibiotic-resistant microorganisms. The discovery of new virulence factors is one of the strategies
44 adopted in pursuing this goal. Along this line, over the past years methionine sulfoxide reductase (Msr) enzymes have
45 gained significance as contributors of virulence for several bacterial strains. Msrs perform the reconversion of methionine
46 sulfoxide to methionine in proteins, and are classified on the basis of their stereoselectivity toward the two
47 diastereoisomers of methionine sulfoxide: MsrA isoforms reduce methionine-*S*-sulfoxide, whereas MsrB isoforms reduce
48 methionine-*R*-sulfoxide [8]. The ubiquitous distribution of Msrs, from prokaryotes to eukaryotes, highlights the strategic
49 role they play against oxidative stress, by repairing the oxidative damage inflicted to sensitive protein-bound methionines,
50 and by participating in a cyclic oxidation/reduction mechanism in which methionines, free or bound in proteins, act as
51 scavengers of oxidants [8]. Several studies shown that MsrA enzymes play a role in the virulence of *Staphylococcus*
52 *aureus*, *Salmonella typhimurium*, *Streptococcus gordonii*, *Mycobacterium smegmatis* and *Mycobacterium genitalium* [9].
53 Furthermore, evidence suggested that MsrA could be involved in the transition of *Staphylococcus epidermidis* from
54 commensalism to pathogenicity [9]. MsrA knock-out strains of these microorganisms showed reduced virulence with
55 respect to wild-type strains, in properties such as the ability to survive inside phagocytic cells, the defense against
56 oxidative attack by neutrophils, the colonization of host tissue, and cytotoxicity and adhesion to host cells [9]. MsrB
57 enzymes do not confer significant contribution to virulence in these bacterial strains. Conversely, in *Francisella tularensis*
58 MsrB, but not MsrA, appears to be a key determinant for virulence [9]. Furthermore, in *Pseudomonas aeruginosa*,
59 *Enterococcus faecalis*, *Streptococcus pneumoniae*, *Helicobacter pylori* and *Escherichia coli* both MsrA and MsrB
60 enzymes are engaged in the promotion of virulence, in the resistance to phagocytosis by macrophages and in contrasting
61 the oxidative insult by neutrophils [9]. Finally, upregulation of *msrA* gene in *Streptococcus aureus* appears to occur in
62 response to cell wall-active antibiotics, indicating a possible role of MsrA in antibiotic resistance [10].

63 3. CONCLUSION

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65 While most literature proposes MsrA as a very important virulence factor in some bacterial strains, little is still known
66 about MsrB. More detailed studies are needed to understand the exact function of these intriguing proteins and their
67 mechanism of regulation in prokaryotes. A deeper insight into these aspects could help in stimulating the development of
68 innovative and effective antimicrobial therapies in the field of gene therapy, an alternative solution for the management of
69 a wide range of infectious diseases that are not amenable to standard clinical approaches. This could circumvent the
70 planetary plague of antibiotic resistance and mitigate the deleterious effects of climate change on human health, in view of
71 the ineluctable further increase of global mean temperature by the end of this century and that could be more severe
72 respect the optimistic scenario prefigured by the Paris Climate Agreement in 2015 [11].

74 COMPETING INTERESTS

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76 Authors have declared that no competing interests exist.

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