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CsrA-controlled proteins reveal new dimensions of Acinetobacter

baumannii desiccation tolerance

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1 ABSTRACT

2 Hospital environments are excellent reservoirs for the opportunistic pathogen Acinetobacter baumannii in part because it is exceptionally tolerant to desiccation. We found that relative to 3 4 other A. baumannii strains, the virulent strain AB5075 was strikingly desiccation resistant at 2% 5 relative humidity (RH), suggesting that it's a good model for studies of the functional basis of 6 this trait. Consistent with results from other A. baumannii strains at 30% RH, we found the 7 global post-transcriptional regulator CsrA to be critically important for desiccation tolerance of 8 AB5075 at 2% RH. To identify CsrA-controlled proteins that may contribute to desiccation 9 tolerance we used proteomics to identify proteins that were differentially present in wild type and csrA mutant cells. Subsequent mutant analysis revealed nine genes that were required for wild 10 type levels of desiccation tolerance, five of which had modest phenotypes. Catalase and a 11 12 universal stress protein gene were moderately important for desiccation tolerance and two 13 genes of unknown function had very strong desiccation phenotypes. The predicted amino acid 14 sequence of one of these genes predicts an intrinsically disordered protein. This category of 15 proteins is widespread in eukaryotes but less so in prokaryotes. Our results suggest there may 16 be mechanisms responsible for desiccation tolerance that have not previously been explored in 17 bacteria.

18

19 **IMPORTANCE**

Acinetobacter baumannii is commonly found in terrestrial environments but can cause
nosocomial infections in very sick patients. A factor that contributes to the prevalence of *A*. *baumannii* in hospital settings is that it is intrinsically resistant to dry conditions. Here, we
established the virulent strain *A. baumannii* AB5075 as a model for studies of desiccation
tolerance at very low relative humidity. Our results show that this trait depends on two proteins
of unknown function, one of which is predicted to be an intrinsically disordered protein. This
category of protein is critical for the small animals named tardigrades to survive desiccation. Our

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27 results suggest that *A. baumannii* may have novel strategies to survive desiccation that have
28 not previously been seen in bacteria.

29

30 INTRODUCTION

31 Hospital-acquired infections are an important healthcare concern and economic burden (1, 2)32 and environmental persistence plays a critical role in the transmission of bacteria that cause 33 these infections (3-6). One such bacterium is Acinetobacter baumannii, an opportunistic 34 pathogen that infects very sick patients. It is responsible for about 2% of nosocomial infections 35 in the United States and Europe and the frequencies are higher in the rest of the world. A. baumannii is especially problematic because on a global basis, about 45% of isolates are multi-36 drug resistant (7). A factor that contributes to the prevalence of A. baumannii in hospital settings 37 38 is desiccation tolerance. A. baumannii can survive in a desiccated state on inanimate dry 39 surfaces for days to several months (8–10). These surfaces include materials that are often 40 encountered in the hospital, such as polyvinyl chloride, rubber, and stainless steel (11).

41

42 When desiccated, bacteria must respond to diverse stresses that include accumulation of 43 reactive oxygen species, loss of cytoplasmic volume, and loss of cell membrane integrity (12, 44 13). Proteomics analysis of A. baumannii showed that desiccated cells had higher levels of 45 proteins involved in protein stabilization, antimicrobial resistance, and reactive oxygen species 46 detoxification (14). Attributes of A. baumannii, that have been shown to be associated with 47 desiccation tolerance include biofilm formation (15, 16) and protein aggregation (17). Lpx M_{AB} dependent acetylation of lipid A is essential for survival of A. baumannii ATCC17978 at 40% RH 48 (18), and a recA mutant of ATCC17978, defective in DNA repair, had a pleiotropic phenotype, 49 50 including a defect in desiccation tolerance (19). katE, encoding catalase also contributes to 51 desiccation tolerance (20). Despite these observations, the number of genes identified in A. baumannii that are specifically involved in desiccation tolerance is small. This could be because 52

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53 A. baumannii cells have evolved modified cell structures that are both essential for viability and 54 important for desiccation tolerance. The genetic basis for this would be difficult to uncover in mutant screens. It is also possible that the conditions of desiccation used, typically 30% RH in 55 studies to date, were not sufficiently severe to allow identification of some desiccation tolerance 56 57 genes. 58 59 With the goal of identifying new desiccation tolerance genes, we established A. baumannii strain 60 AB5075 as a model for studying desiccation tolerance under severe conditions of 2% RH. We 61 then followed up on a recent report showing that csrA, which encodes a global posttranscriptional regulator, is important for desiccation tolerance (21) and identified nine CsrA-62 controlled proteins that confer desiccation tolerance on AB5075 at 2% RH. One of these has 63 64 predicted properties that suggest new dimensions of desiccation tolerance.

65

66 **RESULTS**

Desiccation assay. Previous studies have shown that *A. baumannii* can survive in a desiccated
state for days to several months (8–11, 20). For these and other desiccation studies,

69 investigators worked with a variety of strains and usually incubated cells at either 30% RH or in 70 room air, which varied between 25 and 61% RH in one study (20). These differences can make 71 it difficult to compare desiccation phenotypes between studies. Thus we thought it important to 72 establish a robust desiccation assay that reduces experimental variables like choice of strains,

drying times, and RH during desiccation, using a highly desiccation resistant strain.

74

Following from previous reports, saturated calcium chloride hexahydrate solution placed in a
sealed plastic Snapware container caused the RH inside the container to rapidly equilibrate to
30% (9). We found that use of DRIERITE instead of calcium chloride, resulted in an RH of 2%.
To test desiccation tolerance under different conditions, we grew bacteria to a desired density in

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79 Trypticase Soy Broth (TSB), harvested them, washed them twice, and resuspended them in 80 phosphate buffer to a final OD₆₀₀ of 1. Drops of cell suspension were placed on polycarbonate membranes and filtered to allow for rapid drying. The membranes were then placed in 81 82 uncapped 15 ml conical centrifuge tubes and incubated in desiccation containers. After various 83 periods of incubation, buffer was added to each centrifuge tube followed by 5 min of shaking on a rotary shaker. Viable cell numbers were then determined by plating on TSB agar. To control 84 for the stress of filtration we did viable counts immediately following filtration and took this as our 85 "day 0" time point. 86

87

Relative desiccation tolerance of A. baumannii strains. As shown in Fig 1A, A. baumannii 88 strain AB5075 and Escherichia coli strain W3100 each survived desiccation at 30% RH far 89 90 better than Pseudomonas aeruginosa strain PAO1. However at 2% RH, A. baumannii survived 91 far better than either E. coli or P. aeruginosa. As has been reported (20, 22), we found that A. 92 baumannii stationary phase cells were much more tolerant to desiccation than actively growing 93 cells (Fig. S1) and so we routinely used stationary phase cells in our desiccation assays. We 94 compared the desiccaton tolerance of AB5075 to two additional frequently used laboratory 95 strains of A. baumannii, ATCC17978 and ATCC19606. We found that AB5075 was strikingly more resistant than the others at 2% RH and somewhat mores resistant at 30% (Fig. 1B). 96 97 Given the strength of its phenotype, these findings indicate that AB5075 is a good model for 98 studing desiccation tolerance. This strain was isolated from a surgical wound, is multidrug 99 resistant and is highly virulent in an animal model (23). A comprehensive ordered mutant library of AB5075 is available that has two to three segenced Tn insertions in each gene and is called 100 101 the three-allelle library (24).

102

CsrA is critical for desiccation tolerance. We examined the contribution of the post transcriptional regulator CsrA to desiccation tolerance of AB5075 by constructing a *csrA*

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105	deletion mutant ($\Delta csrA$). We found that the mutant grew poorly on TY agar and had an
106	elongated cell morphology when grown in TY broth (Fig. 2A). On agar plates, large colonies
107	frequently appeared on a backgound of poor growth, likely due to occurance of second site
108	suppressor mutations in the $\Delta csrA$ strain. The $\Delta csrA$ strain was also defective in growth on other
109	nutrient-rich media, including LB broth, nutrient broth, and TSB broth. A similar sensitivity to
110	growth in complex media was reported by Farrow et al. for several A. baumannii strains
111	including strain AB5075 (21). In agreement with Farrow et al., a $\Delta csrA$ mutant grew as the wild
112	type in defined medium, in our case, M9 minimal medium with 10 mM succinate as a sole
113	carbon source (M9/succinate), and it had close to a wild type cell morphology (Fig 2A). A
114	Yersinia enterocolitica csrA mutant, has a growth defect in LB due to the presence of 90 mM of
115	NaCl (25). However, the A. baumannii $\triangle csrA$ mutant was not sensitive to this level of NaCl. In
116	fact, the mutant grew in M9/succinate supplemented with up to 100 mM of NaCl without a
117	significant reduction of growth compared to the wild type.
118	
119	When desiccated after growth in M9/succinate to stationary phase, the AB5075 <i>AcsrA</i> mutant

lost almost all viability over 6 days (Fig. 2B and Table 1). The desiccation phenotype was complemented by expressing *csrA in trans*. $\Delta csrA$ mutant cells incubated for 6 days after being filtered and resuspended in PBS remained fully viable (Fig 2B).

123

124 **CsrA affects mutiple cellular processes in** *A. baumannii*. To identify genes whose 125 translation might be regulated by CsrA, we compared the proteomes of wild-type and $\Delta csrA$ 126 cells (Table S2A). There were 97 proteins present at higher levels in the $\Delta csrA$ mutant 127 compared to the wild type (ratio of $\Delta csrA/WT \ge 2.5$, Table S2B). Among these were proteins for 128 type IV pilus assembly, synthesis of the siderophore ferric acinetobactin, and a 129 glutamate/aspartate transporter. The $\Delta csrA$ mutant also had elevated levels of enzymes for the

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130	catabolism of hydroxcinnamates, phenylacetate and quinate. Levels of an alcohol
131	dehydrogenase (ABUW_1621) and an aldehyde dehydrogenase (ABUW_1624) were also
132	elevated. The $\Delta csrA$ mutant was defective in pilus-mediated twiching motility as assessed by
133	movement aross a soft-agar plate (Fig. 3A). The mutant also had a severe growth defect when
134	grown on succinate in the presence of ethanol (Fig. 3B). One possible explanation for this is that
135	the csrA mutant metabolized ethanol to form toxic acetaldehyde to levels that slowed growth.
136	
137	There were 106 proteins present in lower amounts in the $\Delta csrA$ mutant compared to the wild
138	type (ratio of WT/ $\Delta csrA \ge 2.5$, Table S2C). A large proportion of these (39%) are annotated as
139	hypothetical proteins. Several membrane proteins, and proteins annotated as involved in eta -
140	lactam antibiotic resistance (ABUW_1194, 2619, and 3497), trehalose synthesis (ABUW_3123)
141	and possibly biofilm formation (ABUW_0916) were in lower abundance in the $\Delta csrA$ mutant
142	compared to wild type. As reported previously, a $\Delta csrA$ mutant did not form biofilms (21). This
143	phenotype was complemented by expressing the <i>csrA</i> gene <i>in trans</i> (Fig. 3C). The \triangle <i>csrA</i>
144	proteome profile also suggested that CsrA is invovled in promoting the expression of proteins
145	invovled in oxidative stress, including peroxidase (ABUW_0628) and catalase (katE,
146	ABUW_2436). Indeed the $\Delta csrA$ mutant lacked detectable catalase activity (Fig. 3D).
147	
148	Genes important for desiccation tolerance in A. baumannii AB5075. We took advantabge
149	of the three-allele transposon library to test how important some of the gene transcripts that
150	were likely to be controlled by CsrA were for desiccation tolerance (Tables 1 and S1). When
151	possible, we tested two different transposon mutants (transposon insertions in different

positions of the gene) for each gene. *A. baumannii* AB5075 produces opaque and translucent

153 colony variants that interconvert at high frequency and reflect changes in the thickness of

154 capsular exopolysaccharide (26). AB5075 cells with decreased capsule production are about

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2.5-fold more sensitive to desiccation (27). Here, we used only opaque colonies of AB5075 andits mutant derivatives in our desiccation assays.

157

We found that katE and ABUW 2639 mutants were about 5-fold more sensitive to desiccation 158 159 than the wild type, whereas ABUW 2433 and ABUW 2437 mutants were greater than 100-fold more sensitive to desiccation than the wild type (Table 1, Fig 4). The phentoytpes of 160 161 ABUW 2433 and ABUW 2437 mutants could be complemented (Fig S2). The ABUW 2433 162 protein has 411 amino acids and is annotated as a KGG domain-containing protein. The KGG 163 domain comprises a small region in the N-terminus of the protein and the remainder of the protein is annotated by InterPro as a disordered region that includes a series of AT hook DNA 164 binding motifs (SMART SM00384). The full length ABUW 2433 sequence was predicted to be 165 166 intrinsically unstructured when queried with the IUPred3 tool (https://iupred.elte.hu) (28). 167 ABUW 2437 is annotated as an iron-containing redox enzyme or a heme-oxygenase -like 168 protein (Fig 4). The predicted ABUW 2437 transcript has traits characterisitic of a target of CsrA 169 post-transcriptional regulation. The DNA sequence predicts a relatively long 5' untranslated 170 region (316 bp) and there is a predicted CsrA binding motif (GGA) in the ribosome binding site 171 of the transcript. ABUW 2639 is annoated as belonging to a universal stress protein A family. It 172 has been shown to protect A. baumannii ATCC17987 from oxidative stress of hydrogen 173 peroxide (29).

174

We wondered if *ABUW_2433* and *ABUW_2437* might play a role in promoting desiccation
tolerance of the two *A. baumannii* strains, ATCC17978 and ATCC19606, that do not survive
well at 2% RH (Fig 1B). ATCC19606 has the gene region shown in Fig 4 intact, but the gene
that is homologous to *ABUW_2433*, encoding the KGG domain-containing protein, is annotated
as a pseudogene. ATCC17978 appears to be missing a gene in homologus to *ABUW_2433*.
However it has conitguous *katE* and iron-containing redox protein genes (*A1S 1386 and*

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A1S_1385). Expression of the two AB5075 genes *in trans* improved the survival of the two
 ATCC strains at 2% RH (Fig 5), providiing evidence that ABUW_2433 and ABUW_2437 are
 generally important for desiccaton tolerance.

184

185 Other possible desiccation tolerance genes. As shown in Table 1, we identified an 186 additional six genes that are likely regulated by CsrA, that may have a small role in desiccation 187 tolerance. ABUW 0916, encoding a biofilm-associated protein and otsA, encoding tehalose-6-188 phosphate synthase, are the only two of the six for which we can hypothesize some connection 189 to desiccation. Biofilms have been shown to be important for desiccation tolerance of bacteria (15, 16) and trehalose has been shown to play a significant role in desiccation tolerance of 190 eukaryotes and bacteria (30, 31). When added extrinsically to cultures, trehalose increased the 191 192 desiccation tolerance of A. baumannii ATCC 19606 (22). However, a AmtID-otsB mutant of 193 ATCC19606, defective in endogenous production of the compatible solutes, mannitol, and 194 trehalose, was not more sensitive to desiccation than the wild type (22).

195

196 **DISCUSSION**

Depletion of water during desiccation leads to loss of membrane integrity and accompanying 197 198 disruption of aerobic respiration results in the generation of reactive oxygen species, including 199 hydrogen peroxide (32). So, it makes sense that *katE*, encoding catalase, contributes to 200 desiccation tolerance. Proteomics analyses of A. baumannii showed that proteins involved in 201 redox defense including catalase, alkyl peroxidase reductases and superoxide dismutase were 202 elevated in stationary-phase cells (33), which is consistent with the observation made by many 203 that cells stationary-phase cells survive desiccation much better than exponentially growing 204 cells.

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206 Since the desiccation-tolerance genes ABUW 2433 and ABUW 2437 are near or adjacent to 207 katE, on the genome and all are likely to be controlled by CsrA, it seemed important to consider that they might somehow mediate oxidative stress tolerance even though the amino acid 208 209 sequences of the encoded proteins don't have motifs typically associated with reactive oxygen 210 species detoxification. However, we were unable to demonstrate that ABUW 2433::Tn and 211 ABUW 2437:: Tn mutants were sensitive to hydrogen peroxide and paraguat - both powerful 212 oxidizing agents. In addition, a study that looked at effects of hydrogen peroxide exposure on 213 gene expression in A. baumannii, found that katE but not ABUW 2433 or ABUW 2437, was 214 expressed at elevated levels and neither of these genes is part of the OxyR regulon that controls the response to oxidative stress in A. baumannii (34). Thus, we do not think that either 215 216 of these proteins, which have the greatest defects in desiccation found to date, are likely to 217 function by protecting cells against oxidative stress 218

219 The predicted physical properties of ABUW 2433 provide suggestions as to how it may 220 function. It is an intrinsically disordered protein (IDP) that is highly hydrophilic, with 27% 221 positively charged amino acids residues and 31% negatively charged residues. It is also 222 predicted to assume a collapsed or extended conformation, likely depending on its context (ROBETTA PFRMAT TS prediction; https://robetta.bakerlab.org). ABUW 2433 has 13 repeated 223 224 AT-hook DNA binding motifs that extend across about 70% of the protein. This motif 225 preferentially binds to AT-rich sequences in the minor groove of DNA. AT-hook DNA binding 226 motifs are found primarily in eukaryotic proteins, many of which have roles in transcriptional regulation (35, 36). Only 8.5% of annotated AT hook DNA binding motifs are found in bacteria, 227 but about half of these are found in gamma proteobacteria, the group to which A. baumannii 228 229 belongs. We hypothesize that ABUW 2433 binds to A. baumannii DNA and somehow protects 230 it from desiccation-induced damage. IDPs are critical for the microscopic animals called tardigrades to survive desiccation. When desiccated, some of these proteins vitrify and probably 231

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trap desiccation sensitive molecules in a noncrystalline amorphous matrix, thereby protecting
them from denaturation or other forms of destruction (37, 38). IDPs or proteins with IDP
domains are less common in prokaryotes than in eukaryotes, but drawing from work on
eukaryotes, they are proposed to play a central role in cellular process in bacteria that may
depend on the formation of molecular condensates (39). It is possible that this is important for
the viability of desiccated *A. baumannii*.

238

Although not much work has been done on Acinetobacter CsrA, based on what is known for 239 240 other gamma proteobacteria, we hypothesize that a set of ncRNAs that is induced by a GacSA (ABUW 3306 and ABUW 3639) two-component regulatory system, controls the repressor 241 activity of CsrA by sequestering it (40). We can draw a link between the GasSA system and 242 243 CsrA because they both appear to control catabolism of the aromatic compound phenylacetate. An A. baumannii AgacA mutant is unable to catabolize phenylacetate (41), and our proteomics 244 results suggest that CsrA acts to repress the synthesis of at least one enzyme required for 245 246 phenylacetate degradation. In fact, we have determined that the $\Delta csrA$ mutant grows better on 247 phenylacetate; with a doubling time of 39 min, than the wild type, which has doubling time of 50 248 min. We hypothesize that a $\Delta gacA$ mutant does not synthesize ncRNAs that would normally "sponge-up" CsrA, thus allowing CsrA to bind to the 13 -gene (ABUW 2524 to ABUW 2536) 249 phenylacetate mRNA transcript to repress its translation. At this point we do not have a clear 250 251 understanding of the inventory of A. baumannii ncRNAs that may bind to CsrA, but ncRNAs are 252 abundant in AB5075, and several of them are expressed at extremely high levels (42). The 253 desiccation phenotype of CsrA appears to depend on its ability to activate translation and 254 although it's difficult to reconcile this activity with a model where CsrA is sequestered by ncRNAs, it is known that ncRNA turnover can occur resulting in the release of free CsrA (43). 255 256 Most of what is known about mechanisms of CsrA action centers on its role as a repressor of

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translation (44–47) and it may be of interest to probe its capability as an activator in *A*. *baumannii*.

259

We found that A. baumannii AB5075 survived desiccation for six days at 2% RH much better 260 261 than two other A. baumannii strains that we tested, but it is important to note that most studies 262 of desiccation tolerance have been carried out at about 30% RH or in room air and the 263 emphasis has been on the number of days or months that a particular strain remains viable 264 when desiccated. When Farrow et al (20) tested the survival of several strains that were dried 265 and incubated at an RH of 25–61% (mean 46%) they found AB5075 to have an average survival time of 90 days, whereas strains ATCC19606 and ATCC17978 had average survival 266 times of 3 and 34 days respectively. Even though AB5075 is tolerant to desiccation over 267 268 months at a mean RH of 46% and over days at 2% RH, we cannot necessarily conclude that the 269 same sets of genes are needed for desiccation tolerance under these two different conditions. 270 For example, Farrow et al. (20) found that the response-regulator protein BmfR was important 271 for desiccation tolerance of ATCC17978 in long term desiccation assays, whereas we did not 272 observe a role of *bmfR* in protecting AB5075 from desiccation in shorter term incubations at 273 2%RH (Table S1).

274

275 Here we established a robust assay for desiccation tolerance of a highly virulent strain of A. baumannii and identified two genes, ABUW 2433 and ABUW 2437, that are extremely 276 277 important for desiccation tolerance. Outside of the Acinetobacter genus, ABUW 2437 has homologs in Pseudomonas stutzeri (40% amino acid identity). However, except for a partial 278 279 homolog found in Enterobacteriaceae bacterium TzEc051 (99% identity over 38% of the 280 AB5075 protein), ABUW 2433 does not appear outside Acinetobacter. This and the unusual 281 predicted physical properties of ABUW 2433 as anIDP, suggest that there is novel physiological basis for desiccation tolerance in A. baumannii that remains to be explored. 282

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284 MATERIALS AND METHODS

Bacterial strains and growth conditions. Strains used in this study are listed in Table S3A. Strain AB5075 (AB5075-UW) was used as a wild type (24) and individual transposon mutants were obtained from the Manoil lab comprehensive ordered transposon mutant library at the University of Washington (24). All strains except for the $\Delta csrA$ mutant were routinely grown and maintained in TY (10 g Tryptone, 5 g Yeast extract, and 8 g NaCl in 1000 ml) medium or BBL Trypticase Soy Broth (TSB) media at 37°C, unless otherwise stated. The $\Delta csrA$ mutant was grown in M9/succinate.

292

293 Desiccation assay. Strains from a frozen stock (-80°C) were streaked onto TY plates and 294 incubated at 37°C. Colonies (three to five) were picked and inoculated into 2 ml of TSB, and 295 cultures were grown overnight at 37°C with a shaking speed of 200rpm. Overnight cultures were 296 diluted to yield an initial OD₆₀₀ of 0.025 in 10 ml TSB in a 50 ml Erlenmeyer flask. Cultures were grown at 37°C with a shaking speed of 200 rpm to mid-exponential-phase (OD₆₀₀=0.4 to 0.6) or 297 stationary-phase (24 hours after inoculation). Cells were harvested by centrifugation and 298 washed twice with Dulbecco's phosphate-buffered saline (DPBS, Gibco), and cell density was 299 adjusted to OD_{600} =1 (about 5 x 10⁸ cells/ml) with DPBS. Cell suspension (2 spots of 50 µl each 300 301 per membrane) was filtered onto a 0.4 µm Whatman nucleopore polycarbonate track-etched membranes (25 mm diameter) that had been placed in Nalgene analytical filter units, and the 302 303 membranes were then placed into 15 ml uncapped centrifuge tubes. To obtain the T0 (baseline) viable cell number, 1 ml of DPBS was immediately added to one tube and incubated for 5 min at 304 room temperature $(24 \pm 2^{\circ}C)$ on a rotary shaker. Viable cell numbers were determined by 305 plating on TSB agar. For desiccation, tubes with membranes were placed in a Snapware 306 containers (2.3 x 6.3 x 8.4 inches) that contained DRIERITE in the lids of 50 ml centrifuge tubes 307 308 (x4, 7.5 g of DRIERITE desiccant in each lid) or saturated calcium chloride hexahydrate solution

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309	in 5 ml beaker (x8) to yield the RHs of 2% or 30% (\pm 2), respectively. The Snapware containers
310	were incubated at room temperature. Digital hygrometers (VWR International Ltd) were placed
311	in each container to monitor the RH. At desired time points, tubes containing membranes were
312	removed from the containers, 1 ml of DPBS was added to each tube, and incubated for 5 min at
313	room temperature on a rotary shaker. Viable cell counts were determined on TSB agar. For
314	each strain, a minimum of three biological replicates of desiccation assays were performed
315	except for individual transposon mutants, which were assayed twice for each allele.

316

Construction of the \triangle *csrA* **mutant**. In-frame deletion of the *csrA* (*ABUW* 2750) gene was 317 318 generated by overlap extension PCR as described (48). PCR primers are listed in Table S3B. 319 PCR product was cloned into mobilizable suicide vector pEX2-TetRA and transformed into E. 320 coli NEB 10-beta (New England Bio Labs). The sequence-verified deletion construct was transformed into E. coli S17-1, and further mobilized into A. baumannii strain AB5075 by 321 conjugation on TY agar. Single recombinant conjugants were first selected on M9/succinate 322 plate containing 20 µg/ml Tc, and Tc resistant colonies were further plated onto M9/succinate 323 324 plate containing 5% sucrose. Sucrose resistant and Tc sensitive colonies were screened by 325 colony PCR and sequencing to validate the expected chromosomal in-frame deletion of the csrA 326 gene.

327

To complement the $\Delta csrA$ mutant, the full length *csrA* gene plus the 15 bp upstream that contains the putative ribosome binding site was PCR amplified and cloned into pMMB67EH-TetRA. The construct was transformed into *E. coli* NEB 10-beta (New England Bio Labs). The sequence-verified construct was transformed into *E. coli* S17-1, and further mobilized into the $\Delta csrA$ mutant by conjugation on M9/succinate plates containing 20 µg/ml Tc. As a negative control, empty vector pMMB67EH-TetRA was used. The same procedure was used to clone $ABUW_2433$ and $ABUW_2437$ for complementation experiments.

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336 **Phenotypic characterization of the** \triangle *csrA* **mutant.** M9/succinate was used in all experiments. For motility assays, an overnight culture (16 to 18 hours) was diluted to yield $OD_{600}=0.5$, and 2 337 338 µl of sample was spotted onto the freshly prepared M9/succinate plate and incubated at 37°C 339 for 24 h. For biofilm assays, an overnight culture was inoculated into 100 µl of M9/succinate in 340 Costar vinyl 96 well "U" bottom plates (initial $OD_{600}=0.05$), and the plates were sealed with 341 Breath-Easy sealing membranes. After incubation at room temperature for 48 h, culture was 342 removed, the plate was rinsed with tap water twice, and 150 µl of 0.1% crystal violet solution 343 was added to each well. After incubating at room temperature for 15 min, crystal violet solution was removed, the plate was rinsed with tap water 5 times, and the plate was dried at room 344 temperature. For catalase assays, cells were harvested at OD₆₀₀=0.5, supernatant was 345 346 removed, and cells were resuspended in DPBS to yield 10 mg wet cell/100 µl DPBS. 100 µl of 347 cell suspensions were placed in 13 x 100 mm borosilicate glass tubes. Then 100 µl of 1% Triton 348 X-100 and 100 µ1 of 30% hydrogen peroxide were added, mixed thoroughly, and incubated for 349 15 min at room temperature (49).

350

351 **Label-free protein quantification.** Since the $\triangle csrA$ mutant had a severe growth defect on TY 352 medium, both the wild type and $\Delta csrA$ mutant were grown in M9/succinate. Two biological sample 353 replicates were prepared for each strain. Cells from each culture were harvested at OD₆₀₀=0.5 by centrifugation, washed twice with DPBS, and cells were stored in -80°C before further analysis. 354 355 Cells were lysed in buffer containing 4% SDS, 100 mM Tris pH8.0, 10 mM DTT by heating at 95°C for 5 min. After cooling to room temperature, the lysates were sonicated with ultrasonication 356 probe on ice to shear DNA. Total protein concentration was determined by the BCA assay 357 358 (Thermo Pierce, Rockford, IL). 500 µg of each protein lysate was reduced and alkylated, diluted 359 in 8 M urea solution, and the SDS was removed with a 3kD molecular weight cutoff filter. After buffer exchange, the protein lysates were digested with trypsin (Promega, Madison, WI) at 37°C 360

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361 overnight and the digested samples were desalted with 1cc C18 Sep-Pak solid phase extraction 362 cartridges (Milford, MA, Waters). The eluted samples were vacuum dried and resuspended in 0.1% formic acid. Reverse phase nanoLC-MS analysis of the protein digests was carried out with 363 364 a Thermo Easy-nanoLC coupled to a Thermo Q-Exactive Plus Orbitrap mass spectrometer. 365 Triplicate top 20 data-dependent acquisition runs were acquired for each sample, and 1 µg of protein digest was loaded for each run. The peptides were separated by a 50 cm x 75 µm I.D C8 366 367 column (5 µm diameter, 100 Å pore size C8 MichromMagic beads) with a 90 min 10 to 30% B aradient (solvent A: 0.1% formic acid in water, solvent B: 0.1% formic acid in acetonitrile, flow rate 368 300 nl/min). The MS data acquisition parameters were set as follows: full MS scan resolution 70k, 369 maximum ion injection time 100 mS, AGC target 10⁶, scan range of 400 to 2000 m/z; MS/MS scan 370 resolution 17.5 k. maximum ion injection time 100 mS. AGC target 5⁴, isolation window 1.6 m/z. 371 HCD NCE 35 scan range of 200 to 2000 m/z; loop count 20, intensity threshold 5³, underfill ratio 372 1%, dynamic exclusion 10 sec. High resolution MS^2 spectra were searched against a target-decov 373 374 proteome database of strain AB5075 (a total of 7678 sequences) downloaded from Uniprot (Oct17, 2017) using Comet (version 2015.02 rev. 1) (50) with following parameters: precursor 375 peptide mass tolerance 20 ppm, allowing for -1, 0, +1, +2, or +3 ¹³C offsets; fragment ion mass 376 377 tolerance 0.02 Da; static modification, carbamidomethylation of cysteine (57.0215 Da); variable 378 modification, methionine oxidation (15.9949 Da). The search results were further processed by 379 PeptideProphet (51) for probability assignment to each peptide-spectrum match, and ProteinProphet (52) for protein inference and protein probability modeling. The output pepXML 380 381 files from three technical replicates were grouped for subsequent spectral counting analysis using Abacus (53). The pepXML and protXML files for each sample, combined ProteinProphet file from 382 all samples were parsed into Abacus for spectral counting of each protein. The following filters 383 384 were applied for extracting spectral counts from MS/MS datasets: (1) the minimum 385 PeptideProphet score the best peptide match of a protein must have maxIniProbTH=0.99; (2) The 386 minimum PeptideProphet score a peptide must have to be even considered by Abacus,

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iniProbTH=0.50; (3) The minimum ProteinProphet score a protein group must have in the COMBINED file, minCombinedFilePw=0.90. Spectral counts for 1616 proteins were reported across four sample groups (two strains and two biological replicates) with estimated protein false discovery rate of 1.94%. The protein expression fold changes between wild type AB5075 and $\Delta csrA$ mutant were computed from adjusted spectral counts output from Abacus. The mass spectometry data have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository with the dataset identifier xxxxxxxx.

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395

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399 us to the possibility that intrinsically disordered proteins could be involved in desiccation

400 tolerance in *A. baumannii*.

401

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409 Table 1. Genes that contribute to desiccation tolerance of *A. baumannii* AB5075.

Gene	Gene name	Gene annotation	Relative viability loss ratio (day 6/day 0 desiccation)ª	Protein abundance (ratio of WT/ <i>∆csrA</i>)
AB5075 (WT)			1.0	
ABUW_0916		Biofilm-associated protein	1.5	2.6
ABUW_2433		KGG domain-containing protein	125	4.3
ABUW_2436	katE	Catalase	7.5	3.2
ABUW_2437		Heme oxygenase-like protein	150	10.5
ABUW_2639		Universal stress protein family	6.0	4.4
ABUW_2724		Hypothetical protein	4.0	2.4
ABUW_2750	csrA	Carbon storage regulator	480	8.5
ABUW_3123	otsA	Trehalose-6-phosphate synthase	2.0	10.5
ABUW_3346	acnA	Aconitate hydratase 1	1.5	3.2

410 ^a All mutants were except the ∆csrA strain were grown in TSB broth to the stationary phase of growth

prior to being desiccated. The viability loss of wt day6/day 0 was 17 for the WT grown in TSB broth. The

412 viability loss of WT grown in M9/succinate was 276

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415 **FIGURE LEGENDS**

Figure 1. Desiccation tolerance of *A. baumannii*. (A) Comparison of AB5075, *E. coli* W3110, and *P. aeruginosa* PAO1 at 30% RH or 2% RH. (B) Desiccation tolerance of *A. baumannii* strains AB5075, ATCC17978, and ATCC19606 after 0 days (control) and 6 days of desiccation at 30% RH or 2% RH. The cell numbers represent the total number of viable cells recovered from each membrane. The data are the average of three or more biological replicates, and standard deviations are shown as error bars.

422

423 **Figure 2.** Growth and desiccation tolerance of a $\triangle csrA$ mutant (A) Comparison of wild type (WT)

424 and a $\triangle csrA$ mutant on TY and M9/succinate agar and in TY and M9/succinate broth. (B)

425 Desiccation tolerance of WT, $\triangle csrA$ mutant, $\triangle csrA$ mutant with pMMB (empty vector), and $\triangle csrA$

mutant with *csrA* expressed in trans (pAB2750) at day 0 and at day 6 of desiccation at 2% RH
or at 6 days in PBS. The cell number represents the total number of viable cells recovered from

428 each membrane. The data are the average of three or more biological replicates, and standard429 deviations are shown as error bars.

430

Figure 3. Phenotypes of the $\Delta csrA$ mutant: (A) twitching motility; (B) growth of wild type (black symbol) or the $\Delta csrA$ mutant (red symbol) in M9/succinate in the absence (closed symbol) or presence (open symbol) of 0.5% ethanol. The data shown are representative of each strain and condition; (C) crystal violet staining of biofilms from wild type, $\Delta csrA$ mutant, $\Delta csrA$ mutant with pMMB (empty vector), or $\Delta csrA$ mutant with pAB2750; and (D) catalase activity of wild type, $\Delta csrA$ mutant, $\Delta csrA$ mutant with pMMB (empty vector), or $\Delta csrA$ mutant with pAB2750.

438 **Figure 4.** *A. baumannii* genes important for desiccation tolerance. (A) Genes and (B)

439 Desiccation tolerance of mutants at 2% RH. The cell number represents the total number of

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- viable cells recovered from each membrane. The data are the average of three or more
- 441 biological replicates, and standard deviations are shown as error bars.

442

- 443 **Figure 5.** Effect of expression of *ABUW_2433* (red bars) or *ABUW_2337* (blue bars) gene in
- 444 *trans* on desiccation tolerance of *A. baumannii* (A) ATCC17978 and (B) ATCC19606 after 0
- days (control) and 6 days of desiccation at 2% RH. Empty vector (pMMB, black bars) was used
- 446 as a control. The data are the average of three or more biological replicates, and standard
- 447 deviations are shown as error bars.

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SUPPLEMENTAL MATERIAL

Figure S1. Desiccation tolerance of *A. baumannii* AB5075 cells from the log- and stationaryphases of growth. The cell number represents the total number of viable cells recovered from each membrane. The data are the averages of three or more biological replicates, and standard deviations are shown as error bars.

Figure S2. Complementation of desiccation phenotypes of (A) *ABUW_2433*::Tn and (B) *ABUW_2437*::Tn after 0 days (control) and 6 days of desiccation at 2% RH. Empty vector (pMMB) was used as a control. The data are the average of three or more biological replicates, and standard deviations are shown as error bars.

Table S1. Mutants from the three-allele library tested for desiccation tolerance

Table S2. (A) Label-free protein quantification of wild type and the $\triangle csrA$ mutant. (B) List of proteins up-regulated (ratio of $\triangle csrA/WT \ge 2.5$ and $\triangle csrA$ read count ≥ 7.5) in the $\triangle csrA$ mutant compared to wild type. (C) List of proteins down-regulated (ratio of $\triangle csrA/WT \le 0.4$ and WT read count ≥ 7.5) in the $\triangle csrA$ mutant compared to wild type.

Table S3. (A) Bacterial strains used in this study. (B) Plasmids and primers used in this study

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Figure 1. Desiccation tolerance of *A. baumannii*. (A) Comparison of AB5075, *E. coli* W3110, and *P. aeruginosa* PAO1 at 30% RH or 2% RH. (B) Desiccation tolerance of *A. baumannii* strains AB5075, ATCC17978, and ATCC19606 after 0 days (control) and 6 days of desiccation at 30% RH or 2% RH. The cell numbers represent the total number of viable cells recovered from each membrane. The data are the average of three or more biological replicates, and standard deviations are shown as the error bars.



Figure 2. Growth and desiccation tolerance of the $\triangle csrA$ mutant (A) Comparison of wild type (WT) and the $\triangle csrA$ mutant on TY and M9/succinate agar and in TY and M9/succinate broth. (B) Desiccation tolerance of WT, $\triangle csrA$ mutant, $\triangle csrA$ mutant with pMMB (empty vector), and $\triangle csrA$ mutant with csrA expressed in trans (pAB2750) at day 0 and at day 6 of desiccation at 2% RH or at 6 days in PBS. The cell number represents the total number of viable cells recovered from each membrane. The data are the average of three or more biological replicates, and standard deviations are shown as the error bar.

WT csrAMB 150

W^NACSIAMB 150) ACSIA (PAB2150) ACSIA (PAB2150)

10²



Figure 3. Phenotypes of the $\triangle csrA$ mutant: (A) twitching motility; (B) growth of wild type (black symbol) or the $\triangle csrA$ mutant (red symbol) in M9/succinate in the absence (closed symbol) or presence (open symbol) of 0.5% ethanol. The data shown are the representative of each strain and condition; (C) crystal violet staining of biofilms from wild type, $\triangle csrA$ mutant, $\triangle csrA$ mutant with pMMB (empty vector), or $\triangle csrA$ mutant with pAB2750; and (D) catalase activity of wild type, $\triangle csrA$ mutant, $\triangle csrA$ mutant with pMMB (empty vector), or $\triangle csrA$ mutant with pAB2750; and (D) catalase activity of wild type, $\triangle csrA$ mutant, $\triangle csrA$ mutant with pMMB (empty vector), or $\triangle csrA$ mutant with pAB2750.



Figure. 4. *A. baumannii* genes important for desiccation tolerance. (A) Genes (B) Desiccation tolerance of mutants at 2% RH. The cell number represents the total number of viable cells recovered from each membrane. The data are the average of three or more biological replicates, and standard deviations are shown as the error bar.



Figure 5. Effect of expression of ABUW_2433 (red bar) or ABUW_2337 (blue bar) gene *in trans* desiccation tolerance *A. baumannii* (A) ATCC17978 and (B) ATCC19606 after 0 days (control) and 6 days of desiccation at 2% RH. Empty vector (pMMB, black bar) was used as a control. The data are the average of three or more biological replicates, and standard deviations are shown as the error bar.



Supplementary Figure 1. Desiccation tolerance of *A. baumannii* AB5075 cells from the log-phase and stationery-phase of growth. The cell number represents a total number of viable cells recovered from each membrane. The data are the averages of three or more biological replicates, and standard deviations are shown as error bars.



Supplementary Figure 2. Complementation of desiccation phenotypes of (A) ABUW_2433::Tn and (B) ABUW_2437::Tn after 0 days (control) and 6 days of desiccation at 2% RH. Empty vector (pMMB) was used as a control. The data are the average of three or more biological replicates, and standard deviations are shown as the error bars.