



# **What Happens in the Staphylococcal Nucleoid under Oxidative Stress?**

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**Abstract:** The evolutionary success of *Staphylococcus aureus* as an opportunistic human pathogen is largely attributed to its prominent abilities to cope with a variety of stresses and host bactericidal factors. Reactive oxygen species are important weapons in the host arsenal that inactivate phagocytosed pathogens, but *S. aureus* can survive in phagosomes and escape from phagocytic cells to establish infections. Molecular genetic analyses combined with atomic force microscopy have revealed that the MrgA protein (part of the Dps family of proteins) is induced specifically in response to oxidative stress and converts the nucleoid from the fibrous to the clogged state. This review collates a series of evidences on the staphylococcal nucleoid dynamics under oxidative stress, which is functionally and physically distinct from compacted *Escherichia coli* nucleoid under stationary phase. In addition, potential new roles of nucleoid clogging in the staphylococcal life cycle will be proposed.

**Keywords:** *Staphylococcus aureus; Escherichia coli;* oxidative stress; nucleoid; MrgA; Dps; nucleoid associated protein; atomic force microscopy

### 1. Introduction

The application of atomic force microscopy (AFM), providing direct observation of bacterial nucleoids, has given informative clues that are followed by critical findings in the molecular mechanisms and physiology of prokaryotic systems [1,2]. Nucleoids that are experimentally dispersed from lysed cells are usually observed as fibrous structures in both Gram-positive and Gram-negative bacteria, and also in organelles of prokaryotic origins [3,4]. However, the components of the nucleoids are diverse among bacterial species and their amounts and constituent components undergo dynamic changes depending upon environmental conditions. Such dynamic behavior of the nucleoid components can be linked to the transition of physical characteristics of the nucleoid. This short review summarizes what is known about the staphylococcal nucleoid, especially focusing on its unique morphological change under oxidative stress, and discusses its potential relevance in the life of this important human pathogen.

### 2. Staphylococcus aureus Lifestyle and Importance of Oxidative Stress Resistance

*Staphylococcus* belongs to the Gram-positive *Bacilli* class of Firmicutes that contains a low G/C content in the genome. This class also includes *Bacillus* and *Listeria* spp. The genus *Staphylococcus* is composed of about 60 species [5], and the most clinically relevant one is *S. aureus. S. aureus* asymptomatically inhabits our nasal cavity but is a major opportunistic pathogen responsible for a

broad spectrum of infections ranging from superficial skin abscesses to more severe life-threatening diseases such as pneumonia, sepsis, and toxic shock syndrome. Staphylococcal infections, both in hospitals and in the community, are serious problems in clinical settings, largely because of the difficulty in antibiotic treatment arising due to acquired resistance [6].

*S. aureus* has to cope with a variety of environmental stresses and bactericidal factors in host environments [7]. These include desiccation, hyperosmolarity [8–10], and the immune system [11,12]. Once *S. aureus* is phagocytosed, it is challenged by a series of host bactericidal factors such as acidic pH, antimicrobial peptides, and reactive oxygen species (ROSs). A series of host enzymes and "Fenton reaction" are responsible for the generation of ROSs. NADH oxidase generates superoxide anion from oxygen [13] while superoxide dismutase (SOD) catalyzes its conversion into hydrogen peroxide [14,15]. Ferrous iron (Fe<sup>2+</sup>) then converts the hydrogen peroxide into the highly reactive hydroxyl radical (this process is called the Fenton reaction) [16,17].

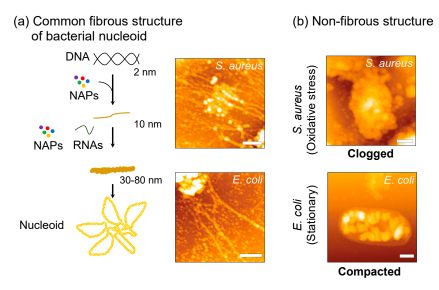
*S. aureus* can survive in professional phagocytes such as neutrophils and macrophages [18–20], where the staphylococcal antioxidant enzymes responsible for the detoxification of ROSs must play critical roles. The anti-oxidant enzymes include SOD [21,22], catalase (that converts hydrogen peroxide into H<sub>2</sub>O and O<sub>2</sub> [23]), and the metallo regulon gene A (MrgA) [24]. It is considered that the ability to survive in migratory phagocytes allows *S. aureus* to spread within our body and to induce severe recurrence or chronic infection [19]. This process relies on multiple regulatory factors, such as Agr (quorum sensing accessory gene regulator) and SigB (general stress response sigma factor), but not SarA (global regulator) [19]. Sortase A and virulence factors such as alpha-toxin, aureolysin, protein A, are also involved in this process [19]. Thus, fine-tuning of the relevant gene expression is necessary in the initial and the following phases of the infection.

#### 3. Oxidative Stress Induces Nucleoid Clogging

#### 3.1. Characteristics of S. aureus Nucleoid in Comparison with Other Bacteria

Most bacterial genomes are circular. In bacteria, genomic DNA (in the scale of a few cm) is packed in a cell (with a diameter of a few  $\mu$ m) in the form of the "nucleoid" with a variety of proteins, RNAs, low-molecular weight compounds, etc. [25]. In contrast to the interphase eukaryotic genome that is separated by the nuclear envelope from the cytosol, the prokaryotic genome is established in the cytosol without a nuclear envelope; i.e., the prokaryotic genome function is achieved in harmony with replication, transcription, and translation all occurring in the cytosolic environment. A variety of methods to isolate the nucleoid have demonstrated different aspects of nucleoid structures, nucleoid-associated proteins (NAPs), role of RNA, and low molecular weight compounds, etc. For example, electron microscopy observations of the nucleoid isolated under high salt conditions have revealed that the circular fibrous genome in bacteria, as a whole, is bundled in the core portion and forms a rosette-like structure with interwound loops emanating radially from the core [26–28].

The nucleoid released from cells lysed under physiological salt concentrations is observed as a fibrous structure with variable thickness regardless of the bacterial species: *S. aureus, Escherichia coli,* and *Clostridium perfringens* [3] (Figure 1a). The fiber thickness ranges between 30~80 nm in width with NAPs and RNAs as structural components [29]. Treatment of the released nucleoids by RNase A, which digests mainly single-stranded RNA [30], makes the nucleoid fibers narrow down to 10 nm, but never releases the naked DNA (2 nm) [29]. In addition, neither RNase III nor RNase H can release the 10-nm fibers. Treatment with rifampicin that targets RNA polymerase to prohibit the transcription also increases the proportion of 10-nm fibers. Thus, nascent RNAs and single-stranded RNAs are involved in the 30~80 nm fibrous nucleoid. It is likely that RNAs are interwoven to thicker fibers in the released nucleoid structure. These hierarchical organizations seem to be general characteristics of bacterial nucleoid. RNase treatment also converts thick nucleoid fibers to thinner ones in the organelles of prokaryotic origin; i.e., chloroplasts and mitochondria [4].



**Figure 1.** Nucleoid structure and dynamics in bacteria. (a) A model of the structural hierarchy of bacterial nucleoid proposed by a series of dissection analyses with AFM and on-substrate lysis method [2]. Naked DNA (2 nm thickness) is complexed with nucleoid-associated proteins (NAPs) to form fibers 10 nm in width, which are a fundamental structural unit to form thicker fibers as well as the compacted nucleoid [31]. Scale bar: 500 nm. (b) Non-fibrous structures. Staphylococcal nucleoid is clogged under oxidative stress, but not in the stationary phase. In contrast, *E. coli* nucleoid is compacted in the stationary phase. Scale bar: 500 nm. Original source of AFM images is [7].

Protease treatment of *E. coli* nucleoid releases not only 10-nm fibers but also naked DNA [29], suggesting that NAPs are structurally important components in nucleoid organization. The E. coli nucleoids isolated under mild salt concentrations with spermidine consist of a set of DNA-binding proteins including the RNA polymerase subunits and about 300 species of transcription factors [32,33]. Among them, Hu (heat-unstable nucleoid protein), HNS (histone-like nucleoid structuring protein), IHF (integration host factor protein), StpA (suppressor of T4 td mutant phenotype A, H-NS homolog), Dps (DNA-binding protein from starved cells), Fis (factor for inversion stimulation), and Hfq (host factor for phage RNA Q $\beta$  replication) were historically believed to be the major nucleoid proteins that were structurally and functionally important [34]. Hfq is now recognized as an RNA chaperone that governs post-transcriptional regulation [35], although another role of Hfq has been implicated in plasmid replication, transposition, and transcription [36]. On the other hand, Hfq was shown to alter the DNA topology indirectly rather than directly associating with DNA [37]. Here it is interesting to note that the released nucleoid from lysed cells of single deletion mutant strains of E. coli (i.e., deletion mutants of genes encoding Hu (hupA, hupB), HNS (hns), IHF (himA, himD), StpA (stpA), Fis (fis), and Hfq (*hfq*)) sustained the fiber structure of 10 nm [29]. This result suggests that each protein is not essential by itself to build up 10-nm fibers.

These proteins are shared among Gram-negative bacteria. However, other than Hu and Dps homologues, they are missing in the genomes of Gram-positive bacteria including *S. aureus* [2]. Namely, irrespective of the structural similarity of the nucleoids mentioned above, many NAPs are diverse depending on the species [38]. *S. aureus* has an Hfq homologue with a substantial RNA binding activity [39]. However, its function still remains elusive since its deletion exhibits no phenotypic changes [40].

Subtractive proteomic analysis of the nucleoid isolated under physiological salt concentrations with spermidine identified staphylococcal proteins that exclusively exist in the nucleoid fraction, but not in soluble cytosol and membrane fractions. They were termed csNAPs (contamination subtracted list of NAPs). The complete lists of 92 csNAPs-log (log phase), 141 csNAPs-st (stationary phase), and 113 csNAPs-ox (oxidative stress) are available in [38,41]. The top 50 csNAPs, sorted by the emPAI values that reflect the protein abundance, are summarized in Table 1. Staphylococcal

csNAPs contains global regulators, fatty acid synthesis enzymes, oxidoreductases, and ribosomal proteins [41], which are common features in bacterial nucleoids [38], and is reasonable if we consider the environmental differences between prokaryotic and eukaryotic genomes (i.e., the absence and presence of nuclear envelope).

		Oxidative Stress	
ID	Gene	Annotation	emPAI
sau:SA1414	rpsT	30S ribosomal protein S20	3.45
sau:SAS033	rpmF	50S ribosomal protein L32	3.34
sau:SA0092		hypothetical protein	3.23
sau:SA2032	rplR	50S ribosomal protein L18	2.72
sau:SA1663		UPF0342 protein SA1663	1.82
sau:SA0093		hypothetical protein	1.55
sau:SA1504	infC	Translation initiation factor IF-3	1
sau:SA2043	rpsS	30S ribosomal protein S19	0.89
sau:SA1074	fabG	3-oxoacyl-[acyl-carrier-protein] reductase	0.72
sau:SA1279	gpsB	Cell cycle protein gpsB	0.69
sau:SA2022	rplQ	50S ribosomal protein L17	0.65
sau:SA2062	sarV	HTH-type transcriptional regulator sarV	0.64
sau:SA1404	rpsU	30S ribosomal protein S21	0.62
sau:SA0133	dra	Deoxyribose-phosphate aldolase	0.59
sau:SA0162	aldA	Putative aldehyde dehydrogenase aldA	0.58
sau:SA0957		UPF0637 protein SA0957	0.56
sau:SA1053	rpoZ	DNA-directed RNA polymerase subunit omega	0.51
sau:SA0204	acpD	FMN-dependent NADH-azoreductase	0.49
sau:SA0232	lctE	L-lactate dehydrogenase 1	0.47
sau:SA0307	nanE	Putative N-acetylmannosamine-6-phosphate 2-epimerase	0.43
sau:SA1305	hu	DNA-binding protein HU	0.42
sau:SA0365	ahpF	Alkyl hydroperoxide reductase subunit F	0.42
sau:SA1922	rpmE2	50S ribosomal protein L31 type B	0.41
sau:SA0366	ahpC	Alkyl hydroperoxide reductase subunit C	0.4
sau:SA0367	nfrA	NADPH-dependent oxidoreductase	0.4
sau:SA1081	rpsP	30S ribosomal protein S16	0.39
sau:SA1471	rpmA	50S ribosomal protein L27	0.39
sau:SA1116	rpsO	30S ribosomal protein S15	0.38
sau:SA2036	rplX	50S ribosomal protein L24	0.34
sau:SA0468	hprT	Hypoxanthine-guanine phosphoribosyltransferase	0.34
sau:SA0478	<i>pdxT</i>	Glutamine amidotransferase subunit pdxT	0.32
sau:SA0488	, syc	Cysteinyl-tRNA synthetase	0.32
sau:SA0573	sarA	Transcriptional regulator sarA	0.26
sau:SA2029	rplO	50S ribosomal protein L15	0.25
sau:SA2423	clfB	Clumping factor B	0.25
sau:SA1901	fabZ	(3R)-hydroxymyristoyl-[acyl-carrier-protein] dehydratase	0.24
sau:SA0512	ilvE	Probable branched-chain-amino-acid aminotransferase	0.24
sau:SA0520	sdrD	Serine-aspartate repeat-containing protein D	0.24
sau:SA0480	ctsR	Transcriptional regulator ctsR	0.22
sau:SA1172	guaC	GMP reductase	0.22
sau:SA0537	thiD	Phosphomethylpyrimidine kinase	0.22
sau:SA0544		UPF0447 protein MW0542; heme peroxidase	0.22
sau:SA1583	rot	HTH-type transcriptional regulator rot	0.2
sau:SA0772	Y772	UPF0337 protein SA0772	0.2
sau:SA0818	rocD	Ornithine aminotransferase 2	0.2
sau:SA0977	isdA	Iron-regulated surface determinant protein A	0.2
sau:SA0942	def	Peptide deformylase	0.19
sau:SA1032	sepF	Cell division protein sepF	0.19
sau:SA1468	ruvA	Holliday junction ATP-dependent DNA helicase ruvA	0.10
sau:SA2046	rplD	50S ribosomal protein L4	0.17
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Table 1. 50 csNAPs sorted by emPAI values in each condition.

## Table 1. Cont.

		Log Phase		
ID	Gene Annotation			
sau:SA0944 phdB		Pyruvate dehydrogenase E1 component subunit beta	2.39	
sau:SA1414	rpsT	30S ribosomal protein S20	2.06	
sau:SA2033	rplF	50S ribosomal protein L6	1.91	
sau:SA0723	clpP	ATP-dependent Clp protease proteolytic subunit	1.67	
sau:SA0504	rpsG	30S ribosomal protein S7	1.2	
sau:SA1382	sodA	Superoxide dismutase [Mn/Fe] 1	1.18	
sau:SA0729	tpi	Triosephosphate isomerase	1.18	
sau:SA1663	1	UPF0342 protein SA1663	1.17	
sau:SA0366	ahpC	Alkyl hydroperoxide reductase subunit C	0.95	
sau:SA0456	spoVG	Putative septation protein spoVG	0.83	
sau:SA2036	rplX	50S ribosomal protein L24	0.81	
sau:SA1073	fabD	Malonyl CoA-acyl carrier protein transacylase	0.7	
sau:SA1930	rpoE	Probable DNA-directed RNA polymerase subunit delta	0.66	
sau:SA1113	rbfA	Ribosome-binding factor A	0.66	
sau:SA2312	ddh	D-lactate dehydrogenase	0.63	
sau:SA1404	rpsU	30S ribosomal protein S21	0.62	
sau:SA0856	spxA	Regulatory protein spx	0.56	
sau:SA0050 sau:SA2029	rplO	50S ribosomal protein L15	0.56	
sau:SA2029 sau:SA1901	fabZ			
sau:SA1901 sau:SA0719	-	(3R)-hydroxymyristoyl-[acyl-carrier-protein] dehydratase Thioredoxin reductase	0.53 0.53	
sau:SA0719 sau:SA2039	trxB			
	rpmC	50S ribosomal protein L29	0.51	
sau:SA2026	infA . D	Translation initiation factor IF-1	0.49	
sau:SA0245	ispD	2-C-methyl-D-erythritol 4-phosphate cytidylyltransferase 2	0.49	
sau:SA0918	purC	Phosphoribosylaminoimidazole-succinocarboxamide synthase	0.49	
sau:SA0941	D	UPF0356 protein SA0941	0.46	
sau:SA0354	rpsR	30S ribosomal protein S18	0.43	
sau:SA1653 sau:SA1305	traP hu	Signal transduction protein TRAP DNA-binding protein HU	0.43 0.42	
sau:SA1359	EF-P	Elongation factor P	0.42	
sau:SA0942		Peptide deformylase	0.41	
sau:SAS074	pdf1	UPF0457 protein SA1975	0.41	
sau:SA1081	une D	30S ribosomal protein S16	0.4	
sau:SA1081 sau:SA2043	rpsP			
	rpsS	30S ribosomal protein S19	0.38	
sau:SA2399		Fructose-bisphosphate aldolase class 1	0.38	
sau:SA0707	1) /	Uncharacterized protein SAB0704	0.37	
sau:SA0128	sodM	Superoxide dismutase [Mn/Fe] 2	0.36	
sau:SA1717	gatC	tRNA(Asn/Gln) amidotransferase subunit C	0.36	
sau:SA0352	rpsF	30S ribosomal protein S6	0.34	
sau:SA0855	trpS	Tryptophanyl-tRNA synthetase	0.34	
sau:SA0437		UPF0133 protein SAB0428	0.34	
sau:SA2427	arcB	Ornithine carbamoyltransferase catabolic	0.33	
sau:SA2127	rpi	Ribose-5-phosphate isomerase A	0.32	
sau:SA1074	fabG	3-oxoacyl-[acyl-carrier-protein] reductase	0.31	
sau:SA0160		Heme-degrading monooxygenase isdI	0.31	
sau:SA2037	rplN	50S ribosomal protein L14	0.3	
sau:SA2089	sarR	HTH-type transcriptional regulator sarR	0.29	
sau:SA2022	rplQ	50S ribosomal protein L17	0.29	
sau:SA0473	folB	Dihydroneopterin aldolase	0.29	
sau:SA0108	sarH1	HTH-type transcriptional regulator sarS	0.27	
sau:SA0573	sarA	Transcriptional regulator sarA	0.26	

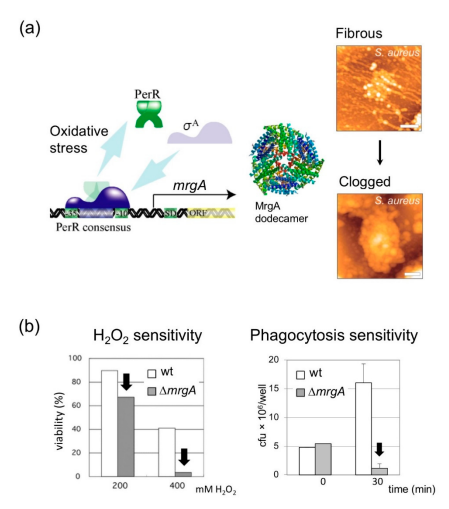
## Table 1. Cont.

		Stationary Phase		
ID	Gene Annotation			
sau:SA0992	trxA	Thioredoxin	4.91	
sau:SA0295		30 kDa neutral phosphatase (Fragment)	3.92	
sau:SA0873		UPF0477 protein SA0873	2.57	
sau:SA1178		UPF0154 protein SSP1415	2.14	
sau:SA1305	hu	DNA-binding protein HU	1.85	
sau:SA1663		UPF0342 protein SA1663	1.82	
sau:SA2043	rpsS	30S ribosomal protein S19	1.6	
sau:SA1067	rpmB	50S ribosomal protein L28	1.55	
sau:SA0456	spoVG	Putative septation protein spoVG	1.47	
sau:SA1909	, atpF	ATP synthase subunit b	1.46	
sau:SA1709	ftn	Ferritin	1.45	
sau:SA0160	isdI	Heme-degrading monooxygenase isdI	1.24	
sau:SA2062	sarV	HTH-type transcriptional regulator sarV	1.1	
sau:SA0760		Glycine cleavage system H protein	1.09	
sau:SA0108	sarH1	HTH-type transcriptional regulator sarH1	1.04	
sau:SAS078	rpmJ	50S ribosomal protein L36	1.01	
sau:SA1904	atpC	ATP synthase epsilon chain	1.01	
sau:SA0032	bleO	Bleomycin resistance protein	0.99	
sau:SA0494	nusG	Transcription antitermination protein nusG	0.98	
sau:SA0478	pdxT	Glutamine amidotransferase subunit pdxT	0.97	
sau:SA2038	rpsQ	30S ribosomal protein S17	0.94	
sau:SA0245	ispD	2-C-methyl-D-erythritol 4-phosphate cytidylyltransferase 2	0.94	
sau:SA1901	fabZ	(3R)-hydroxymyristoyl-[acyl-carrier-protein] dehydratase	0.9	
sau:SA1256	msrB	Peptide methionine sulfoxide reductase msrB	0.89	
sau:SA0128	sodM	Superoxide dismutase [Mn/Fe] 2	0.85	
sau:SA1019		Uncharacterized N-acetyltransferase SA1019	0.85	
sau:SA0437		UPF0133 protein SAB0428	0.81	
sau:SA1074	fabG	3-oxoacyl-[acyl-carrier-protein] reductase	0.72	
sau:SA2431	isaB	Immunodominant staphylococcal antigen B	0.72	
sau:SA1041	pyrR	Bifunctional protein pyrR	0.7	
sau:SA2089	sarR	HTH-type transcriptional regulator sarR	0.65	
sau:SAP018	arsC	Protein arsC	0.59	
sau:SA2040	rplP	50S ribosomal protein L16	0.54	
sau:SA2266		Uncharacterized oxidoreductase SAR2567	0.54	
sau:SA1529		UPF0173 metal-dependent hydrolase SA1529	0.52	
sau:SA1146	bsaA	Glutathione peroxidase homolog bsaA	0.47	
sau:SA1076	rnc	Ribonuclease 3	0.46	
sau:SA0774	The	Probable ABC transporter ATP-binding protein	0.46	
sau:SA0941		UPF0356 protein SA0941	0.46	
sau:SA1461	apt	Adenine phosphoribosyltransferase	0.44	
sau:SA2392	panB	3-methyl-2-oxobutanoate hydroxymethyltransferase	0.44	
sau:SA1206	femA	Aminoacyltransferase femA	0.44	
sau:SA0354	rpsR	30S ribosomal protein S18	0.43	
sau:SA0934	ptsH	Phosphocarrier protein HPr	0.42	
sau:SA1032	sepF	Cell division protein sepF	0.4	
sau:SA0470	hsp33	33 kDa chaperonin	0.1	
sau:SA1471	rpmA	50S ribosomal protein L27	0.39	
sau:SA1081	rpsP	30S ribosomal protein S16	0.39	
sau:SA0704	1731	UPF0230 protein	0.39	
sau:SA0826	spsB	Signal peptidase IB	0.39	
5au.3A0020	эрэр	orginal peptidase in	0.59	

blue: ribosomal protein; red: Hu; yellow: oxidoreductases; green: global regulator; grey: fatty acid metabolism.

#### 3.2. Apparent Correlation between Nucleoid Clogging and Oxidative Stress

In *S. aureus*, the fibrous structures released from lysed cells diminish under oxidative stress conditions and the nucleoids are observed as clogged forms [42] (Figure 1b). The key factor to cause such clogging was found to be MrgA (similar to Dps family proteins in *E. coli*, see Section 3). The *mrgA* gene does not express its gene product without oxidative stress due to transcription suppression by PerR (Figure 2a). Once PerR senses the oxidative stress, it is released from the *mrgA* promoter and *mrgA* transcription is induced. Owing to this tight regulation, MrgA is specifically expressed under oxidative stress conditions, and reaches c.a. 30,000 molecules (2500 dodecamer) per cell [43]. The deletion mutant of *mrgA* is unable to clog the nucleoid under oxidative stress, while artificial over-expression of MrgA by plasmid, or by mutation in the *perR* suppressor gene, results in the nucleoid clogging even under normal growth conditions without the oxidative stress.



**Figure 2.** MrgA is essential for the oxidative stress response. (a) Left: A model of *mrgA* gene regulation. The *mrgA* gene is among the highly up-regulated genes upon phagocytosis [18]. PerR, the suppressor of *mrgA* transcription, can sense oxidative stress and dissociate from the *mrgA* promoter to release the inhibition. By this regulation, MrgA is specifically expressed under oxidative stress conditions and induces the nucleoid clogging [42]. Center: MrgA forms dodecamer like other Dps family proteins [24]. It lacks known DNA binding regions, and how MrgA binds DNA is not known [43]. Right: AFM images of nucleoid dynamics. Scale bar: 500 nm. (b) The *mrgA* gene is essential in hydrogen peroxide resistance [42], as well as in phagocytosis resistance [24]. These resistances are attributed to the ferroxidase activity of MrgA [24]. A *mrgA* deletion increased the sensitivities to H<sub>2</sub>O<sub>2</sub> (left) and the time-dependent phagocytic killing (right). Error bars at the 30 min time point represent SD (*n* = 3). Images and graph data were reproduced from [7,24,42].

Similar, but physiologically and physically distinct, changes in nucleoid dynamics have been observed in *E. coli* (reviewed in [2]) (Figure 1b), where Dps plays a key role. The expression of Dps in *E. coli* is induced by oxidative stress (as a part of the OxyR regulon) as well as in the stationary phase. Dps is the dominant nucleoid protein in the stationary phase [44], and the nucleoid is tightly compacted [45], limiting the access of DNA binding proteins (except for RNA polymerase [46]). However, Dps expression in the log-phase does not compact the nucleoid because a log-phase dominant nucleoid protein, Fis, prevents the compaction [47]. In contrast to *E. coli*, artificial expression of MrgA by plasmid in *S. aureus* results in clogged nucleoid irrespective of the growth phases. The MrgA-expressing cells are not different in the growth rate from the wild type cells, indicating that nucleoid clogging in response to oxidative stress seems to be a phenomenon specific in *S. aureus*, of which physiological relevance is still open to discussion (see the following sections).

#### 4. Is Nucleoid Clogging Required or Not for the Oxidative Stress Tolerance?

## 4.1. MrgA Is a Bifucntional Molecule with Ferroxidase Activity That Is Essential for Oxidative Stress Resistance

*S. aureus* MrgA is important for oxidative stress resistance like other Dps family proteins [42] (Figure 2b). Dps family proteins usually assemble into dodecamers and exert ferroxidase activity. MrgA also assembles into dodecamers and the structural data is available in Protein Data Bank under the accession number of 2D5K [24]. Several, but not all, of Dps family proteins including *E.coli* Dps [48] and staphylococcal MrgA [24] can bind DNA. Scavenging free iron is important to prevent the Fenton reaction that generates the hydroxyl radical from ferrous iron (Fe<sup>2+</sup>) and hydrogen peroxide [49]. There is a report showing that the ferroxidase activity alone, without the DNA binding activity, can contribute to oxidative stress resistance: *Streptococcus mutans* Dpr (Dps-like peroxide resistance gene, Dps-family protein) that can bind iron but not DNA is critical to cope with oxidative stress [50,51].

In *S. aureus*, when the ferroxidase center of MrgA (Asp56 and Glu60) is mutated, the susceptibility to oxidative stress increases [24]. These mutations do not disrupt dodecamer formation and DNA binding activity. Therefore, it can be concluded that; (1) ferroxidase activity is essential, and; (2) DNA binding activity alone is not important for oxidative stress resistance.

## 4.2. DNA Binding Activity of MrgA Is Dispensable for Hydrogen Peroxide Resistance and Survival in Phagosome, but Not for Nucleoid Clogging

While it became evident that the ferroxidase activity of MrgA is important for oxidative stress resistance in *S. aureus* [24], the relevance of DNA binding of MrgA has still been under question. The first point we addressed was whether or not, in addition to the ferroxidase activity, DNA binding of MrgA is essential for the physical protection of the genomic DNA [43]. One difficulty is that the DNA binding domain of MrgA has not been identified, whereas that of E. coli Dps is known to be in the N-terminal region [43]. Since we have been unable to make specific MrgA variants that lack the DNA binding activity so far, we instead introduced the N-terminal-deletion mutant of *E. coli* Dps ( $\Delta$ 18-Dps) that has no DNA binding activity into the *S. aureus mrgA*-knockout mutant. The obtained results clearly demonstrated that the nucleoid is clogged by the expression of Dps in S. aureus  $\Delta mrgA$ , but not by  $\Delta 18$ -Dps, indicating that DNA binding activity of Dps is necessary for nucleoid clogging. By analogy, MrgA DNA binding activity is likely responsible for the nucleoid clogging. In addition,  $\Delta$ 18-Dps, as well as Dps, compensated for MrgA in hydrogen peroxide resistance regardless of nucleoid clogging, demonstrating that the DNA binding activity is dispensable for such resistance itself. Namely, the molecular mechanisms of DNA clogging and hydrogen peroxide resistance are likely to be independent, although both mechanisms may cross-over, depending upon the environmental conditions.

Furthermore, an interesting implication is that the apparently distinct nucleoid clogging in *S. aureus* and nucleoid condensation in *E. coli* are brought about by similar molecular mechanisms. In other

words, MrgA and Dps can be exchanged in *S. aureus* for nucleoid clogging. This may be a key feature for further investigation of the molecular mechanisms for genome condensation in general.

#### 5. Any Physiological Relevance in Nucleoid Clogging?

#### 5.1. Characteristics of csNAPs in the Clogged and Relaxed Nucleoid

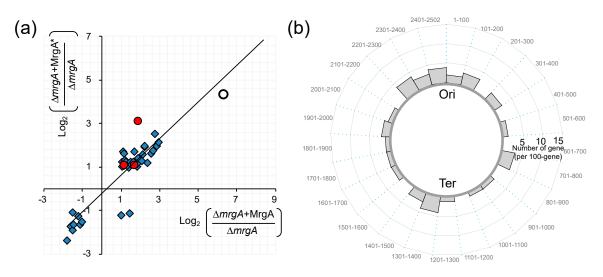
According to the list of csNAPs in nucleoids (Table 1), some specific features in the clogged nucleoid can be extracted. First, Hu, an *E. coli* major NAP, always exists as csNAPs in staphylococcal nucleoid regardless of the growth phases or the presence of oxidative stress. Second, other *E. coli* major NAPs are lost through the evolutionary processes in *S. aureus*. Third, on the other hand, the isolated staphylococcal nucleoid contains so-called global regulators (Sar homologues and Rot). These would be the evolutionary distinct staphylococcal counterparts of the *E. coli* major NAPs.

These global regulators are constitutively expressed components of the nucleoid in any conditions (log, stationary, and oxidative stress). Such steady state expression of *S. aureus* global regulators makes a striking contrast to the drastic exchange of *E. coli* major NAPs from the log (Fis abundant) to the stationary phase (Dps dominant) [52]. Upon oxidative stress, Sar homologues are maintained in the nucleoid, but some up- and down-regulations among the homologues may take place (see Table 1). The enzymes responsible for detoxification of oxidative stress are also constitutively detected as csNAPs, although the molecular species are diverse depending on the conditions.

As mentioned above, the *S. aureus* nucleoid clogged by MrgA is biologically active and allows cell proliferation. In fact, ribosomal proteins are abundant csNAPs in the clogged nucleoid. In clear contrast, the compacted *E. coli* nucleoid has few ribosomal proteins [38]. The dynamics of csNAPs upon *S. aureus* nucleoid clogging seems to be less drastic than those in *E. coli* nucleoid compaction. Thus, considerable parts of the nucleoid function are sustained in the clogged form.

#### 5.2. Effect of Nucleoid Clogging on Transcriptome Profile

Staphylococcal genome is about 2.8 Mbp and contains c.a. 2500 protein-coding genes (c.a. 85% of the genome) [53]. Interestingly, artificial expression of MrgA or MrgA\* (MrgA carrying mutations in the ferroxidase centre at Asp56 and Glu60) by plasmid in the  $\Delta mrgA$  strain can affect the transcriptome profile similarly in the absence of oxidative stress (Ushijima et al., in preparation for submission): There were 41 signals significantly changed (>2 fold or <0.5 fold) by MrgA and MrgA\*, and MrgA and MrgA\* had the same effect for 39 of them (Figure 3). Most of these signals originated from non-coding sequences (Figure 3, diamonds), and only a few from protein coding sequences (Figure 3, red circles). This observation may reflect the differential expression of small RNAs or the difference in the lengths of mRNAs' untranslated regions. It should be noted that the DNA binding activity of MrgA affected the transcriptome without its ferroxidase activity.



**Figure 3.** (a) Gene expression is affected by MrgA as well as MrgA\* similarly in the absence of oxidative stress. Transcriptome data of "*mrgA* deletion mutant ( $\Delta mrgA$ )", "*mrgA* overexpression ( $\Delta mrgA$ +MrgA)", and "*mrgA*\* overexpression ( $\Delta mrgA$ +MrgA\*)" strains grown in the absence of oxidative stress were obtained by a standard procedure by using GeneChip (Affymetrix). X axis: Comparison between " $\Delta mrgA$ +MrgA" and " $\Delta mrgA$ ". Y axis: Comparison between " $\Delta mrgA$ +MrgA\*" and " $\Delta mrgA$ ". Log<sub>2</sub> fold differences of the loci that showed significant differences (i.e., >2 fold or <0.5 fold) in both comparisons were plotted. Red circles: Protein coding sequences (CDSs). Blue diamonds: Non-CDSs. Open circle: *mrgA*. Thus, MrgA dependent nucleoid clogging can affect the expression of RNAs mainly from non-CDSs in the absence of the oxidative stress. This effect is not due to the ferroxidase activity of MrgA, since the MrgA\* overexpression has similar effects to the MrgA overexpression: The correlation coefficient is 0.897. (b) Location of the genes which were affected by both MrgA and MrgA\* in the absence of oxidative stress. The cumulative numbers of the genes (plotted in graph (a)) per 100-gene region are plotted in a circular way. SA numbers in N315 genome are shown outside the circle: 1 = *SA0001 (dnaA)* through *SA2502*.

On the other hand, 112 protein coding genes were up-regulated and 90 were down-regulated under oxidative stress [54]. Under oxidative stress (20  $\mu$ M PQ: Phenanthrenequinone, [55,56]) WT and  $\Delta mrgA$ , which have clogged and fibrous nucleoids respectively, exhibit distinct profiles in their transcriptomes (91 loci > 2-fold, and 87 loci < 0.5-fold) (Ushijima et al., in preparation for submission). In contrast to Figure 3 (in the absence of oxidative stress), many transcripts from coding sequences were differentially accumulated (listed in Tables 2 and 3). An intriguing feature of this list is the location dependency; many of the genes are located around the replication origin (Ori), and few from around the Ter side (Figure 4). Notably, the expression patterns of genes in the Staphylococcal Cassette Chromosome (SCC) that locates near the Ori were largely distinct between WT(+PQ) and  $\Delta mrgA(+PQ)$ . The lists also present the genes for virulence factors (red), nucleic acid metabolism (green), iron metabolism (*sirC*, *SA0120*), transcription regulators (yellow) including three global regulators (staphylococcal accessory regulators, *sarH1*, *sarY*, *sarV*), and bacteriophage holin/anti-holin.

**Table 2.** Genes with higher expression in WT(+PQ) than  $\Delta mrgA(+PQ)$ . Oxidative stress was given by 20  $\mu$ M PQ at 37 °C for 30 min to log phase cells. Transcriptome was analyzed by a standard procedure by using GeneChip (Affymetrix).

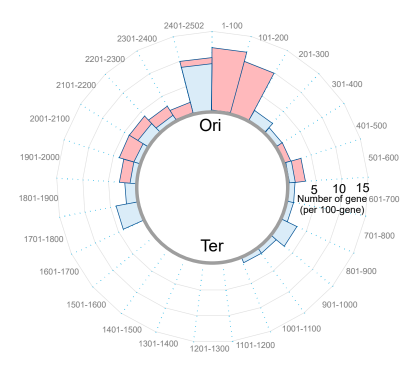
Log-Difference WT(+PQ) /DmrgA(+PQ)	Gene Name	N315 SA Number	Annotation/Similarity
4.04	mrgA(dps)	SA1941	MrgA, Dps family protein
2.07	lrgB	SA0253	antiholin-like protein LrgB
1.86	/	SA2133	hypothetical protein
1.65	оррВ	SA0853	oligopeptide ABC transporter permease
1.59	arcB	SA2427	ornithine carbamoyltransferase
1.54	sen	SA1643	enterotoxin SeN (in pathogenicity island, SaPIn3)
1.53	/	SA2470	histidinol dehydrogenase
1.51	/	SA2417	nisin susceptibility-associated two-component sensor histidine kinase
1.50	/	SA2264	hypothetical protein
1.49	hisZ	SA2472	ATP phosphoribosyltransferase regulatory subunit/ His-tRNA synthase
1.48	/	SA2454	acetyltransferase
1.35	/	SA2429	ArgR family transcriptional regulator
1.31	/	SA0667	7-cyano-7-deazaguanine synthase
1.30	pyrF	SA1047	orotidine 5'-phosphate decarboxylase
1.30	/	SA0846	oligopeptide transport system permease OppC
1.29	/	SA1760	holin-like protein (in phage phiN315)
1.28	/	SA1807	mobile element associated protiein (in phage phiN315)
1.26	/	SA0804	Na+/H+ antiporter family protein
1.22	lrgA	SA0252	murein hydrolase regulator LrgA
1.22	yent1	SA1645	enterotoxin Yent1 (in pathogenicity island, SaPIn3)
1.21	/	SA2469	histidinol-phosphate aminotransferase
1.20	/	SA0582	monovalent cation/H+ antiporter subunit E
1.15	purC	SA0918	phosphoribosylaminoimidazole-succinocarboxamide synthase
1.11	/	SA2189	Ferrochelatase family / cobalamin biosynthesis CbiX/ transcriptional regulator NirR
1.09	ureE	SA2085	urease accessory protein UreE
1.09	/	SA1768	phage tail protein (in phage phiN315)
1.03	/	SA1636	hypothetical protein
1.02	/	SA1675	amino acid ABC transporter permease/substrate-binding protein
1.02	nrdD	SA2410	anaerobic ribonucleoside triphosphate reductase
1.02	ssp	SA0744	secretory extracellular matrix and plasma binding protein
1.01	/	SA0324	mepB family protein
1.00	clfB	SA2423	clumping factor B

blue: holin, anti-holin; red: virulence; yellow: transcription regulator; green: nucleic acid metabolism.

**Table 3.** Genes with lower expression in WT(+PQ) than  $\Delta mrgA(+PQ)$ . Oxidative stress was given by 20  $\mu$ M PQ at 37 °C for 30 min to log phase cells. Transcriptome was analyzed by a standard procedure by using GeneChip (Affymetrix).

Log-Difference WT(+PQ)/DmrgA(+PQ)	Gene Name	N315 SA Number	Annotation/Similarity
-3.03	spa	SA0107	immunoglobulin G binding protein A
-2.41	/	SA0080	membrane protein similar to sulfite exporter TauE/Saff family protein
-2.35	/	SA0100	Na/Pi cotransporter family protein
-2.31	sirC	SA0109	iron compound ABC transporter permease SirC
-1.92	/	SA0090	hypothetical protein
-1.90	sarH1	SA0108	staphylococcal accessory regulator H1
-1.87	lacC	SA1995	tagatose-6-phosphate kinase
-1.84	/	SA0085	hypothetical protein
-1.81	/	SA0061	(in Staphylococcus Cassette Chromosome, SCC)
-1.81	/	SA0077	serine/threonine protein kinase (in Staphylococcus Cassette Chromosome, SCC)
-1.78	lctP	SA0106	L-lactate permease
-1.73	/	SA2092	AraC family transcriptional regulator
-1.68	/	SA0102	myosin-cross-reactive MHC class-II like protein
-1.67	lp18	SA0404	lipoprotein encoded in pathogenicity island (in pathogenicity island, SaPIn2)
-1.63	/	SA0124	capsular polysaccharide biosynthesis glycosyltransfera: TuaA
-1.61	/	SA0120	SbnI, siderophore biosynthesis protein
-1.54	/	SA2230	fmtA-like protein/ beta lactamase
-1.50	/	SA0085	tRNA-dihydrouridine synthase
-1.45	/	SA0099	transmembrane efflux pump protein
-1.43	vraA	SA0533	long chain fatty acid-CoA ligase vraA
-1.42	/	SA2303	ABC transporter permease protein
-1.42	/	SA0097	AraC/XylS family transcriptional regulator
-1.38	sarY	SA2091	staphylococcal accessory regulator Y
-1.21	/	SA0105	hypothetical protein
-1.20	/	SA1826	pathogenicity island protein (in pathogenicity island, SaPIn1)
-1.18	/	SA2274	hypothetical protein
-1.17	/	SA2302	ABC transporter ATP-binding protein
-1.12	/	SA0087	tfoX N-terminal domain protein
-1.08	/	SA0078	hypothetical protein
-1.07	/	SA0037	MaoC domain-containing protein (in Staphylococcus Cassette Chromosome, SCC)
-1.06	/	SA0536	hypothetical protein
-1.05	/	SA2154	hypothetical protein
-1.05	/	SA0088	hypothetical protein
-1.05	<i>lacA</i>	SA1997	galactose-6-phosphate isomerase subunit LacA
-1.04	sarV	SA2062	staphylococcal accessory regulator V
-1.04	hisG	SA2471	ATP phosphoribosyltransferase catalytic subunit
-1.03	/	SAS028	hypothetical protein
-1.00	sodM	SA0128	superoxide dismutase
1.00	fmhA	SA2199	fmhA protein (FemAB like protein,)

red: virulence; yellow: transcription regulator; pale red: iron metabolism; pale blue: oxidative stress related.



**Figure 4.** Location of the genes which are differently expressed between WT(+PQ) and  $\Delta mrgA(+PQ)$  under oxidative conditions. The cumulative numbers of the genes (listed in Tables 2 and 3) per 100-gene region are plotted in a circular way. SA numbers in N315 genome are shown outside the circle: 1 = SA0001(dnaA), through *SA2502*. Genes that were more (blue; Table 2) or less (red; Table 3) expressed in WT(+PQ) than  $\Delta mrgA(+PQ)$  tend to locate around the Ori-side of the genome.

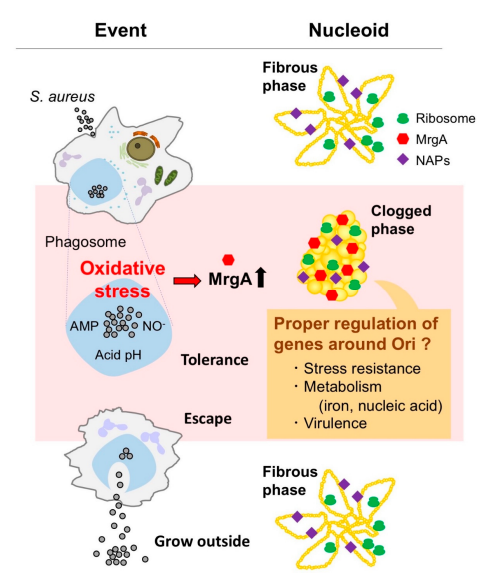
The results from these analyses would define our next strategies towards understanding what is really going on in the cells before and after oxidative stresses. Considerations on the gene regulatory mechanisms before and after nucleoid clogging under oxidative stress conditions will be accelerated, where a series of oxidative stress responsive regulators (such as PerR, MgrA, SarZ, etc) are cooperatively working [57].

In summary: (1) Under no oxidative stress condition, MrgA binding to nucleoid up-regulates specific non-protein coding genes around the whole genome. (2) Under the stress condition, MrgA binding leads to the up-regulation of the protein coding genes near the Ori. (3) Under the stress condition at the same time, MrgA binging down-regulates the protein coding genes near the Ori. These observations clearly lead us to a few interesting implications. First, MrgA binding may cause different nucleoid status with and without oxidative stress. The evidence for this relies on Figures 3 and 4 as well as Tables 2 and 3. Second, one structural or physiological conformation is favored for the expression of non-protein coding genes, and the other is preferred by the up- and down-regulations of specific genes. However, the subtleties of certain distinct gene regulations are not known and left as future questions. Third, most nucleoid functions are supposed to be sustained before and after the nucleoid clogging: In this sense, it is interesting to note that we previously described the 'Armor hypothesis' by postulating the importance of the localization of antioxidant factors in the nucleoid for genome DNA protection [38].

In conclusion, staphylococcal nucleoid is distinct from the well-studied *E. coli* nucleoid in its dynamics of NAP composition and morphologies. Staphylococcal MrgA is specifically expressed under oxidative stress conditions where it plays important roles to support the survival of this opportunistic human pathogen. So far, any role of nucleoid clogging has not been postulated in the oxidative stress resistance. However, it is now clear that the nuclear clogging represents at least two different structural and functional states of the genome; i.e., under physiological oxidative stress and under the

experimental absence of oxidative stress (although whether nucleoid clogging exists physiologically without oxidative stress is still unknown).

A current hypothetical scenario illustrates a certain nucleoid status where gene expression is controlled through the pathogenesis of *S. aureus* (Figure 5). Upon phagocytosis, *S. aureus* senses oxidative stress and induces the expression of MrgA. While the ferroxidase activity directly contributes to the oxidative stress resistance, the DNA binding activity of MrgA converts the nucleoid status into the clogged phase. This may be a preferable state for the proper control of gene expression for survival in phagosomes, as well as preparation for the next step of pathogenesis. Also, it will be an exciting challenge to clarify how particular nucleoid-clogging state is linked to specific gene regulation at the molecular level.



**Figure 5.** Summary and current hypothesis. *S. aureus* is challenged by oxidative stress in phagosomes. The oxidative stress signal is sensed by the PerR transcriptional repressor leading to the induction of MrgA which converts the nucleoid to the clogged state. Unlike condensed *E. coli* nucleoid, the clogged nucleoid sustains the activities of replication and gene expression that are necessary for cell proliferation. Indeed, the clogged nucleoid retains ribosomes and NAPs including the Sar/Rot global regulators. Nucleoid clogging plays no known role in oxidative stress resistance, but it may be that the clogging phase is preferable for the proper expression of the genes locating around the Ori in the genome. Such gene expression is expected to be involved in the tolerance to phagosome-associated stresses, virulence, and prophage activation. It may also affect other nucleoid-related functions.

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