The Human Mitochondrial Genome Encodes for an Interferon-Responsive Host Defense Peptide

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Abstract

The mitochondrial DNA (mtDNA) can trigger immune responses and directly entrap pathogens, but it is not known to encode for active immune factors. The immune system is traditionally thought to be exclusively nuclear-encoded. Here, we report the identification of a mitochondrial-encoded host defense peptide (HDP) that presumably derives from the primordial proto-mitochondrial bacteria. We demonstrate that MOTS-c (mitochondrial open reading frame from the twelve S rRNA type-c) is a mitochondrial-encoded amphipathic and cationic peptide with direct antibacterial and immunomodulatory functions, consistent with the peptide chemistry and functions of known HDPs. MOTS-c targeted *E. coli* and methicillin-resistant *S. aureus* (MRSA), in part, by targeting their membranes using its hydrophobic and cationic domains. In monocytes, IFNγ, LPS, and differentiation signals each induced the expression of endogenous MOTS-c. Notably, MOTS-c translocated to the nucleus to regulate gene expression during monocyte differentiation and programmed them into macrophages with unique transcriptomic signatures related to antigen presentation and IFN signaling. MOTS-c-programmed macrophages exhibited enhanced bacterial clearance and shifted metabolism. Our findings support MOTS-c as a first-in-class mitochondrial-encoded HDP and indicates that our immune system is not only encoded by the nuclear genome, but also by the co-evolved mitochondrial genome.
Introduction

The endosymbiotic theory posits that mitochondria originate from once free-living bacteria that infected eukaryotic ancestral cells. Indeed, owing to their bacterial origin\(^1\), mitochondria still retain several prokaryotic characteristics, including N-formylated proteins and a circular DNA, that register as quasi-self damage-associated molecular patterns (DAMPs)\(^2\). Multiple pattern-recognition receptors (PRRs) can sense DAMPs and elicit pro-inflammatory and type I interferon responses\(^2\). Further, mitochondrial DNA (mtDNA) can be actively released by neutrophils\(^3\) and eosinophils\(^4\) to physically target pathogens and by lymphocytes to signal type I interferon responses\(^5\).

While mtDNA itself can trigger immune responses and directly entrap pathogens, it is not known to encode for genes that yield immune factors. The continuous expansion of our proteome, powered by the characterization of short/small open reading frames (sORFs) in the nuclear\(^6,7\) and mitochondrial\(^8,9\) genomes that yield functional peptides, provides unprecedented opportunities for the discovery of host defense peptides (HDPs), also known as antimicrobial peptides (AMPs)\(^10,11\). All kingdoms of life possess a vast repertoire of AMPs that are now pursued therapeutically as antibiotics, antivirals, cell-penetrating peptides, immunomodulators, and cancer targeting peptides\(^10,12-18\). HDPs are microproteins of 10 to 50 amino acids in length\(^10,12,14\). We have previously identified a mitochondrial-encoded sORF, MOTS-c (mitochondrial ORF from the twelve S rRNA type-c), which yields a 16 amino acid peptide\(^19\). MOTS-c has a key role in regulating cellular homeostasis under cellular stress and during aging\(^20\), in part, by directly translocating to the nucleus to regulate adaptive gene expression\(^19,21-24\).

Based on the endosymbiosis theory, early communication between the proto-mitochondria and proto-eukaryotic cell likely occurred on an immunological basis with immune factors that were encoded within both genomes. Today, bacteria still use gene-encoded peptides, known as bacteriocins, that regulate vital cellular processes (e.g. ribosomal processes, DNA replication, and cell wall synthesis\(^25\)) to control inter- and intra-species proliferation\(^16,26-29\). In the unique endosymbiotic context, these peptides may have served to not only protect the newly formed union from other pathogens, but also to regulate the growth and metabolism of the opposing quasi-self (i.e. the present-day mitochondria and nucleus). Thus, mitochondrial-derived peptides (MDPs)\(^30\), including MOTS-c, may inherently possess immuno-metabolic functions and represent a primordial arm of the eukaryotic immune system (Figure 1A) that evolved with dual roles as cellular regulators during aging\(^19,20,31\).
Here, we report that MOTS-c is a mitochondrial-encoded HDP that directly targets bacteria and regulates monocyte function.

Results

The mitochondrial-derived peptide MOTS-c compromises bacterial viability in vitro

HDPs are characterized by their cationic and amphipathic residues, typically with a net positive charge. Peptide analysis using the hydrophobicity scales of Kyte and Doolittle\(^{32}\) and Sweet and Eisenberg\(^{33}\) indicated that MOTS-c is an amphipathic peptide consisting of a core that is enriched with hydrophobic residues (\(\text{8YIFY}_{11}\) (Figure 1B). MOTS-c retains a +3 charge under physiological pH, largely owing to the basic tail residues (\(\text{13RKLR}_{16}\)) (Figure 1C), which may also serve as a heparin-binding domain (XBBXB-like moiety; B: basic residues) that can bind to cell wall carbohydrates (e.g. glucan, mannan) and mediate bacterial recognition and binding\(^{34,35}\).

Direct bacterial targeting is a hallmark of HDPs\(^{18,36}\). Indeed, MOTS-c dynamically associated with bacteria immediately upon contact (Figures 1D-1H and S1). We mixed MOTS-c (100 µM) with \(E.\ col\)i suspended in water. MOTS-c levels in the media (water) decreased concomitantly with increase detection in cell lysates, indicating direct interaction with bacteria (Figure S1A). HDPs can cause bacterial aggregation, which immobilizes them to the local infection site and enhances pathogen clearance\(^{37-41}\). Consistently, MOTS-c immediately aggregated \(E.\ col\)i and MRSA (methicillin-resistant \(S.\ aureus\)) in a dose- and growth phase-dependent manner (Figures 1D and S1B-1C; Video S1). HDPs engage with bacteria through ionic and hydrophobic interactions owing to their hydrophobic and charged residues\(^{42,43}\). MOTS-c was ineffective in aggregating bacteria under higher salt concentrations (0-1% in ddH\(_2\)O)(Figure 1E), which disrupts ionic interactions and HDP activity\(^{43,44}\). The loss of the hydrophobic and cationic domains of MOTS-c, by substituting the residues with alanine (i.e. \(\text{8YIFY}_{11}>\text{8AAAA}_{11}\) and \(\text{13RKLR}_{16}>\text{13AAAA}_{16}\)), prevented bacterial aggregation (Figures 1F-1G and S1D), consistent with the importance of these residues for HDP function\(^{10,45}\). MOTS-c did not aggregate \(S.\ typhimurium\) or \(P.\ aeruginosa\) (Figure S2), indicating target selectivity independent of Gram status.

HDPs can perturb bacterial membranes via several mechanisms, including progressive blebbing, budding, and pore formation\(^{46-48}\). Using scanning electron microscopy (SEM), we visualized a time-dependent progression of MOTS-c-dependent membrane blebbing in \(E.\ col\)i (Figure 1H) and MRSA (Figure S3). Membrane destabilization by HDPs can also cause bacterial
aggregation\textsuperscript{49-51}. We confirmed rapid compromise of bacterial membrane integrity upon MOTS-c treatment using a fluorescent nucleic acid stain that only penetrates permeabilized membranes\textsuperscript{52} (Figure S4A). Further, MOTS-c treatment depleted cellular ATP levels in \textit{E. coli} (Figure S4B). Consistently, real-time metabolic flux analyses revealed that MOTS-c treatment perturbed respiration and glycolysis (Figure S4C), which requires intact membrane function\textsuperscript{53}.

We next confirmed the antimicrobial effect of MOTS-c on bacterial growth. A single treatment of MOTS-c significantly retarded \textit{E. coli} proliferation in liquid culture in a dose-dependent manner (Figure S4D-E). Two intermediate doses of MOTS-c (50 \textmu M) had comparable effects to a single high-dose treatment (100 \textmu M) (Figure S4D), reflecting the significance of antibiotic treatment frequency in addition to absolute dose. Notably, human HDPs can reach high intracellular concentrations compared to their low circulating levels: LL-37 can reach 40 \textmu M\textsuperscript{54-57} and defensins can constitute 5-7\% of the total protein content of neutrophils\textsuperscript{58,59}. Further, HDPs can reach very high membrane-bound concentrations that are 10,000 times of aqueous solutions (80 mM)\textsuperscript{36}. Interruption of ionic interaction, achieved by higher salt concentration (Figure 1I) or loss of the hydrophobic or cationic domains by alanine-substitution mutagenesis (\textit{i.e.} 8YIFY\textsubscript{11}\textgreater 8AAAA\textsubscript{11} and 13RKLR\textsubscript{16}\textgreater 13AAAA\textsubscript{16}; 100 \textmu M) (Figure 1J), fully reversed the antimicrobial function of MOTS-c on \textit{E. coli} growth. Multiple HDPs also have intracellular roles that contribute to the antimicrobial effect\textsuperscript{60-63}. Using an inducible MOTS-c expression vector, we found that the endogenous overexpression of MOTS-c significantly inhibited \textit{E. coli} (BL21) growth (Figure S4F), indicating a two-stage mechanism of targeting membranes and intracellular components.

\textbf{MOTS-c enhances survival from MRSA exposure in vivo}

To confirm that the antibacterial effects of MOTS-c are sustained \textit{in vivo}, we inoculated mice with MRSA that had been treated with or without MOTS-c (Figure 2A-2C). Whereas the intraperitoneal inoculation of MRSA was lethal, MOTS-c-treated MRSA was not (20\% vs. 100\% survival, respectively) (Figure 2A). However, both groups lost weight initially, but mice inoculated with MOTS-c-treated MRSA started to recover 48 hours post-infection while the surviving population in the control group continued to lose weight (Figure 2B). MOTS-c-treated MRSA, sampled from the inoculated preparation, showed considerably reduced CFU on LB agar plates (Figure 2C). To test whether lethal peritonitis can be induced simply by high levels of PAMPs or required live bacteria, we inoculated mice with heat-killed MRSA and found that they did not cause weight loss nor death (Figure S5). These results suggest that MOTS-c not only opsonizes
bacteria and renders them immunologically unreactive, but also inhibits key pathogenic functions, including proliferation.

*Endogenous MOTS-c is induced upon human monocyte activation in vitro*

Although the initial focus on HDP research was on their direct antimicrobial effect, recent studies have established them as key regulators of immune responses, including monocyte activation and differentiation\(^{10,11}\). Because (i) the discovery of MOTS-c was inspired by a prior study demonstrating interferon gamma (IFNγ)-induced transcripts from the mitochondrial rRNA genes in monocytes\(^{64}\) and (ii) MOTS-c regulates adaptive cellular responses to various types of stress\(^{19,21-24}\), we hypothesized that it may act as a modulator of monocyte activation and differentiation. Endogenous MOTS-c levels dynamically increased in a time-dependent manner upon monocyte-to-macrophage differentiation in (i) primary human peripheral blood monocytes by M-CSF (macrophage colony stimulating factor) (**Figure 3A**) and (ii) a human monocytic cell line (THP-1) by phorbol myristate acetate (PMA)\(^{65}\) (**Figure 3B**). Since they recapitulated the dynamics of MOTS-c upon differentiation signals and they are a tractable cell line, we decided to perform follow-up experiments primarily in the THP-1 system. Endogenous MOTS-c was also induced in THP-1 monocytes in a time-dependent manner following stimulation with lipopolysaccharides (LPS) and IFNγ (**Figure 3C**), a combination of bacterial-derived and cytokine-dependent signals known to synergistically activate monocytes\(^{66-72}\). We then tested if LPS and IFNγ could each induce MOTS-c expression separately. LPS alone increased MOTS-c expression in THP-1 monocytes (**Figure 3D**) and differentiated THP-1 macrophages (**Figure S6A**), consistent with known HDPs\(^{73-81}\). IFNγ alone also induced MOTS-c expression in THP-1 monocytes (**Figure 3E**), consistent with the strong induction of transcripts from the mitochondrial rRNA genes in interferon-induced monocyte-like cells\(^{64}\). Cellular levels of induced MOTS-c in THP-1 macrophages appear to reach high concentrations (**Figure S6B**), consistent with known HDPs\(^{54-59}\).

*MOTS-c regulates the differentiation trajectory of human monocytes in vitro*

Because HDPs can regulate monocyte activation and differentiation\(^{82-85}\), we next tested whether MOTS-c can modulate monocyte differentiation. We previously reported that MOTS-c translocates to the nucleus upon cellular stress to regulate a range of nuclear genes, indicating that mitochondrial-encoded factors can regulate the nuclear genome\(^{21-23,31}\). Indeed, endogenous MOTS-c translocated to the nucleus upon differentiation of THP-1 monocytes by PMA in a time-dependent manner (**Figure 4A**) and following LPS+IFNγ stimulation (**Figure S7A**). Consistent with our previous reports\(^{19,21-24}\), exogenously treated MOTS-c readily entered THP-1 monocytes
(Figure S7B); the uptake was significantly retarded at a lower temperature (4°C) (Figure S7C), indicating the potential involvement of active transport. Exogenously treated MOTS-c peptide also showed strong nuclear localization (Figures 4B and S7D). To test whether MOTS-c regulates nuclear transcriptional programming during early monocyte differentiation, we performed bulk RNA-seq on THP-1 monocytes that were treated with PMA or PMA+MOTS-c for 2 hours (Figure 4C-4H). Multidimensional scaling (MDS) revealed that the overall expression profiles of PMA and PMA+MOTS-c monocytes were distinct, as they were clearly separated in the MDS 2-dimensional space (Figure 4C). Notably, the separation of the two groups occurred mostly on a single dimension (i.e. dimension 1, which captures the largest proportion of variance among samples), compatible with the notion that MOTS-c treatment led to acceleration in differentiation-related gene expression changes. MOTS-c differentially regulated 945 genes between PMA vs. PMA+MOTS-c groups (FDR<5%) (Figure 4D; Table S1), comparable to LL-37 which can affect the expression of >900 nuclear genes in human monocytes.

Using the STRING (Search Tool for the Retrieval of Interacting Genes/Proteins) database version 11.0, which can assess putative changes in protein-protein interaction networks based on our RNA-seq analysis, we identified large gene clusters in the PMA+MOTS-c group, compared to the PMA group, that were related to (i) ribosomes and translation initiation and (ii) chromatin dynamics (Figure 4E; full results in Table S2). Consistently, functional enrichment analysis for Gene Ontology (GO) gene sets revealed that the most significantly targeted functions by MOTS-c included (i) ribosomal and translational processes and (ii) chromatin dynamics (selected terms in Figure 4F; full results in Table S2). We then compared gene expression changes during normal differentiation (control vs. PMA) and MOTS-c-programmed differentiation (PMA vs. PMA+MOTS-c) and Spearman rank correlation (Rho) analysis revealed that changes were significantly correlated, consistent with the notion that MOTS-c treatment can accelerate the normal macrophage differentiation program (Figure 4G). In this context, we identified 64 genes that were specifically attributed to MOTS-c regulation after PMA induction (Figure 4G; Table S1D), of which ribosomal genes were again represented based on STRING analysis (Figure 4H). Consistently, ribosomal levels and differential translation are known to regulate cellular differentiation and lineage commitment, indicating that MOTS-c may broadly impact gene expression during monocyte differentiation.
*In vitro MOTS-c-treated human monocytes yield functionally distinct macrophages*

Due to the impact of MOTS-c on the transcriptome of THP-1 monocytes, we then asked whether MOTS-c can regulate monocyte differentiation to produce macrophages that are functionally distinct. A single exposure to MOTS-c during differentiation enhanced monocyte adherence, an important step for extravasation and differentiation to macrophages. We found that a greater number of MOTS-c-programmed primary human monocytes became adherent 3 days (Figure 5A) and 6 days (Figure S8A) after M-CSF stimulation, indicating an accelerated differentiation program, consistent with our RNA-seq analysis (Figure 4C and 4G). Next, we exposed THP-1 monocytes to MOTS-c at the onset of differentiation and functionally characterized the MOTS-c-programmed macrophages 4 days later. First, MOTS-c-programmed THP-1 macrophages exhibited a significant increase in bacterial killing capacity, determined using a gentamicin protection assay (Figure 5B). Second, we characterized the differential response of MOTS-c-programmed macrophages to LPS stimulation, including cytokine secretion/expression and cellular metabolism (Figure 5C-5F). MOTS-c-programmed THP-1 macrophages exhibited (i) a shift in LPS-induced secretion of IL-1β, IL-1Ra and TNFα as measured by ELISA (Figure 5C) and (ii) selective impact on LPS-induced chemokine and cytokine gene expression as measured by RT-qPCR (Figures 5D and S8B), and (iii) suppressed oxygen consumption in response to LPS, a metabolic reflection of macrophage activity, following acute stimulation (Figure 5E) and later reduced spare respiratory capacity 16 hours post-stimulation (Figure 5F). These results are in line with the previously described impact of LL-37, a well-described human HDP, in reprogramming monocytes to differentiate into macrophages with enhanced antibacterial capacity, suggesting that the mitochondrial-encoded MOTS-c may act in a similar fashion.

*Exposure of primary mouse bone marrow cells to MOTS-c promotes the emergence of a distinct subset of mature macrophages in both sexes throughout aging*

MOTS-c has a significant impact on aging physiology, in part, by acting as a regulator of adaptive stress responses and is considered an emerging mitochondrial hallmark of aging. In addition, aging is also accompanied by maladaptive immune responses, including a shift in macrophage function. Thus, we tested the impact of early MOTS-c exposure on the differentiation trajectories of primary progenitors from the bone marrow of female and male, young and old C57BL/6JNia mice (Figure 6A). Importantly, we used single-cell analyses to investigate heterogeneity in resulting primary macrophage populations (i.e. different macrophage “states”), reflecting their broad spectrum of transcriptional programs trained to dynamically adapt to their environment.
Specifically, we used single-cell RNA-seq (scRNA-seq) to determine whether early exposure (during the first 3 days of differentiation) of primary mouse bone marrow progenitors to MOTS-c may promote the emergence of transcriptionally distinct macrophage subsets. Because of the age-dependent shift in macrophage adaptive capacity\textsuperscript{117}, we collected bone marrow cells from young (4 mo.) and old (20 mo.) mice of both sexes and differentiated them for 7 days +/- MOTS-c into bone marrow-derived macrophages (BMDMs). MOTS-c was given only once concomitantly with M-CSF at the onset of differentiation, and the media was replaced after 3 days in both the control- and MOTS-c-treated conditions (Figure 6A). scRNA-seq was performed at day 7 of differentiation, generating libraries for each biological group separately (N = 5 animals per group, one library for each group obtained by equicellular mixing of BMDMs from each animal). To minimize the negative impact of batch effects, all samples were processed in parallel from bone marrow collection to library preparation and sequencing.

First, we asked whether there were global changes to mature BMDM transcriptomes upon only early exposure to MOTS-c, irrespective of age and sex. For this purpose, we decided to leverage a pseudobulk approach, which best controls false discovery rates\textsuperscript{118}. After aggregating reads for each independent library, we normalized the data using the Variance Stabilizing Transformation from the ‘DESeq2’ R package. Multidimensional scaling (MDS) revealed that macrophages were clearly separated by age on dimension 1 and by MOTS-c treatment on dimension 2 (Figure 6B). Intriguingly, female BMDMs showed greater shift in response to MOTS-c treatment than male BMDMs; notably, female MOTS-c-treated BMDMs appeared “masculinized” (i.e. closer to male samples in MDS space, and with reduced sample-to-sample distance; Figure 6B and S9). With limited sample number, future work will be needed to elucidate interactions of MOTS-c treatment with age and sex in the context of BMDM transcriptional programming. However, our pseudobulk analysis reveals global remodeling of BMDM programs upon transient early exposure to MOTS-c.

Using a shared nearest neighbor (SNN) modularity optimization with the ‘Seurat’ R package, we identified 8 BMDM clusters, indicative of different latent transcriptional states, consistent with a heterogeneous mature macrophage population. Importantly, the 8 distinct single-cell BMDM clusters were comprised of cells from young and old mice of both sexes [cluster labeling is shown on our data using a nonlinear dimensionality-reduction technique, UMAP (uniform manifold approximation and projection)] (Figure 6C). Intriguingly, 2 clusters (clusters 5 and 6) were consistently enriched in MOTS-c-treated condition and their increase was more
pronounced in BMDMs derived from older animals regardless of sex (Figure 6D and 6E; Table S3). To note, aging alone increased the proportion of clusters 5 and 6 (Figure 6D; upper panels), consistent with the connection between MOTS-c signaling and aging19,24,119-121. Cluster 5 was largely enriched in the expression of genes relevant to antigen presentation, whereas cluster 6 showed increased expression of interferon-related genes (Figures 6F-6G and S10-S11; full analysis in Table S3); some genes were shared between these categories. Consistently, overrepresentation analysis using Gene Ontology Biological Process terms (GOBP) showed a clear signature of antigen presentation processes for cluster 5 and IFN-related processes for cluster 6 (FDR < 5%; top 20 pathways in Figure S12; full analysis in Table S4). Notably, interferon signaling can be triggered not only by mtDNA2, but can also induce the expression mtDNA-encoded MOTS-c (Figure 3C and 3E)64. Increased expression of genes involved in antigen presentation and interferon signaling with age has been observed in monocytes and macrophages in mice122-126. In fact, the antigen presentation genes H2-Aa, H2-Ab1, H2-Eb1, CD74, and AW112010 and interferon-related genes Irf7, Ifit2, Ifit3, Ifitm3, and Ifi204 are consistently upregulated with age in our data and that of others (Table S3)122-125, indicating a conserved effect of aging on monocytes/macrophages.
Discussion

Here, we identify MOTS-c as a first-in-class mitochondrial-encoded HDP that can target bacteria and modulate monocyte differentiation. MOTS-c influenced nuclear gene expression during the early phase of monocyte-to-macrophage differentiation to generate distinct macrophage populations that are adapted to bacterial clearance. Our current working model is that MOTS-c initially mounts a preemptive attack on bacteria by aggregating them and preventing growth, then influences monocyte differentiation for efficient microbial clearance. Notably, HDPs possess anti-inflammatory effects upon clearing pathogens to promote a non-inflammatory post-infection resolution and alleviate over-responsive inflammation (e.g. sepsis)\textsuperscript{127,128}, consistent with the systemic anti-inflammatory effects of MOTS-c\textsuperscript{19,21,23,31,129-141}.

Our initial hypothesis was that interbacterial communication using peptides may still exist between bacteria-derived mitochondria and bacteria. Endosymbiosis is a form of sustained infection whereby the primal bacteria and our ancestral cell communicated to coordinate the establishment of mitochondria and eukaryotic life. Such communication was likely derived from immunological processes to regulate each other, which may have laid the foundation for cellular signal transduction. HDPs often regulate highly-conserved cellular functions, such as ribosomes and protein metabolism\textsuperscript{142}, consistent with the regulatory functions of MOTS-c in mammalian cells\textsuperscript{100,101,143-147} (Figure 4E-4H). Evidence for MOTS-c as a therapeutic agent against MRSA in mice adds translational potential for mtDNA-encoded immune peptides for combating bacterial infection\textsuperscript{148}. Further, it is likely that MOTS-c, consistent with other HDPs, may also target viruses. Human myeloblast cells that were infected with Sendai virus to induce IFN expression showed considerable induction of transcripts from the mitochondrial rRNA loci (~75% of induced transcripts)\textsuperscript{64}, a study that influenced the discovery of MOTS-c\textsuperscript{19}. Consistently, we found that MOTS-c is endogenously expressed in monocytes and that it can be induced by IFN\gamma stimulation (Figure 3E).

Our data supports an immunological role for mitochondrial-encoded MOTS-c and provides a proof-of-principle for mitochondrial-encoded HDPs. Together with the ever-increasing identification of sORFs in both our mitonuclear genomes, multiple unannotated HDPs may exist in humans with the potential for clinical use against a broad range of infections\textsuperscript{149,150}. 
Figure Legends

Figure 1. MOTS-c is a mitochondrial-encoded host defense peptide (HDP).
(A) Bacteria and bacteria-derived mitochondria possess gene-encoded immune peptides, known as host defense peptides (HDPs) in higher eukaryotes. (B) MOTS-c has a hydrophobic core (8YIFY11), determined using the hydrophobicity scales of Kyte and Doolittle (K&D)32 and Sweet and Eisenberg (S&E)33. Blue: cationic residues. (C) MOTS-c has a cationic tail (13RKLR16) that confers positive charge (Z) across a pH range. (D) MOTS-c treatment (0-100 µM) immediately aggregates E. coli in a dose dependent manner. (E) MOTS-c-dependent E. coli aggregation is lost in increasing salt concentrations (NaCl; 0-1%) and (F, G) requires its hydrophobic core and cationic domain, consistent with other HDPs. EGFP-expressing E. coli (BL21) shown. WT and mutants devoid of its hydrophobic (8YIFY:8YIFY11>8AAAA11; YIFY) or cationic domain (13RKLR16>13AAAA16; RKLR). Bar, 75 µm. (H) Scanning electron micrographs of E. coli treated with MOTS-c (100 µM) for 0 (immediate fixation), 30, and 60 minutes (n=3). Representative images shown. Bar, 100 nm. (I-J) Growth curve of E. coli (BL21), measured by optical density at 600 nm (OD600), following (n=6) (I) MOTS-c treatment in the presence of 1% NaCl, and (J) treatment with wildtype (WT) MOTS-c and mutants devoid of its hydrophobic (8YIFY11>8AAAA11; YIFY), or cationic domain (13RKLR16>13AAAA16; RKLR). Data expressed as mean +/- SEM. Two-way ANOVA repeated measures. *** P<0.001.

Figure 2. MOTS-c enhances survival from MRSA exposure in vivo.
(A-C) 4x10^8 CFU of mid-log phase MRSA was resuspended in 100 µM MOTS-c or vehicle (water) and immediately injected IP into 16-week-old male C57BL/6J mice (n=4-5). Survival (A) and weight (B) were monitored for 72 hours. (C) 4x10^8 CFU of MRSA resuspended in 100 µM MOTS-c or water was serial diluted and plated on LB agar before injection and colonies counted after overnight incubation. Data expressed as mean +/- SEM. Log-rank (Mantel-Cox) test (A), and Mann-Whitney test (C).

Figure 3. MOTS-c is induced in activated and differentiating monocytes.
(A-B) Total endogenous MOTS-c levels measured as a function of time following monocyte differentiation in (A) primary human monocytes by M-CSF (100 ng/ml) and (B) THP-1 cells by PMA (15 nM). (C-E) Total endogenous MOTS-c levels following THP-1 monocyte activation by LPS (100 ng/ml) and IFNγ (20ng/ml) (C) in combination, (D) LPS alone, and (E) IFNγ alone. Data expressed as mean +/- SEM. Mann-Whitney test. *P<0.05, ** P<0.01, **** P<0.0001.
Figure 4. MOTS-c reprograms early nuclear gene expression during monocyte differentiation.

(A) A time-course measurement of endogenous MOTS-c in purified nuclear extracts following THP-1 monocyte differentiation by PMA (15 nM). (B) Confocal fluorescence images of THP-1 monocytes treated with FITC-MOTS-c (1 µM) for 30 minutes, showing nuclear localization. Nucleus marked by DAPI staining. Bar, 10 µm. (C-H) THP-1 Monocytes were primed with/without MOTS-c (10 µM) for 2 hours, then differentiated with PMA +/- MOTS-c (10 µM) for 2 hours, at which time RNA was collected for bulk RNA-seq analysis (n=6); false discovery rate (FDR) < 5%.

(C) Multidimensional scaling (MDS) analysis across control, PMA, and PMA+MOTS-c groups based on RNA-seq expression profiles after DESeq2 VST normalization. (D) Heatmap of significantly differentially regulated genes by MOTS-c by DESeq2 analysis. (E) Protein-protein interaction network analysis based on genes that were significantly differentially up- and down-regulated by MOTS-c (FDR < 5%) using the STRING (Search Tool for the Retrieval of Interacting Genes/Proteins) database version 11.0. (F) Significantly enriched biological functions based on gene set enrichment analysis (GSEA) using gene ontology (GO). Selected groups are shown, full data in Table S1. (G) Correlation plot of gene expression changes by DESeq2 upon (i) regular induction of differentiation (control vs. PMA) compared to (ii) MOTS-c-directed induction of differentiation (PMA vs. PMA+MOTS-c). Spearman Rank correlation (Rho), and significance of this correlation are reported. Genes that are significantly regulated only upon MOTS-c treatment but not during normal differentiation are highlighted in red and may underlie a specific MOTS-c-induced macrophage state. (H) Protein-protein interaction network analysis based on the 64 genes that were significantly differentially up- and down-regulated by MOTS-c (FDR < 5%) as described in (G) using the STRING database version 11.0. MΦ=macrophage.

Figure 5. MOTS-c promotes the generation of macrophages with enhanced antibacterial capacity. (A) Primary human monocytes were differentiated by M-CSF for 3 days with/without MOTS-c treatment (10 µM; 2-hour priming, then single treatment with M-CSF). Representative images of adhered macrophages (n=6; MΦ=macrophage; Bar, 10 µm). (B-F) THP-1 macrophages were differentiated for 4 days with/without MOTS-c treatment (10 µM; 2-hour priming, then single treatment with PMA). (B) Gentamicin protection assay in MOTS-c-programmed THP-1 macrophages following 1.5- or 3-hours post-infection of E. coli (MOI: 10). CFU: colony forming units (n=6). (C-F) MOTS-c-programmed THP-1 macrophages were stimulated with LPS (100 ng/ml) and (C) secreted levels of IL-1β, IL-1Ra, and TNFα measured by
ELISA after 20 hours (n=6), (D) cytokine expression levels determined by RT-qPCR after 16 hours (n=6), and (E-F) metabolic flux assessed (n=15) by cellular respiration (oxygen consumption rate; OCR) (E) immediately after LPS stimulation and (F) 16 hours after LPS stimulation. Data expressed as mean +/- SEM. Mann-Whitney test, except for (E, F), which used two-way ANOVA repeated measures. *P<0.05, ** P<0.01, *** P<0.001.

Figure 6. MOTS-c generates unique macrophages characterized by enhanced interferon signaling and antigen presentation in an age-related manner.

(A) Single-cell RNA-seq (scRNA-seq) was performed on bone marrow-derived macrophages (BMDMs) from young (4 mo.) and old (20 mo.) mice of both sexes that were differentiated for 7 days in the presence/absence of MOTS-c (10 µM), treated once concomitantly with first exposure to M-CSF, and present only for the first 3 days of differentiation. (B) Multidimensional scaling (MDS) analysis across each of the 8 groups based on pseudobulk gene expression profiles for each biological group after performing DESeq2 VST normalization. (C-D) Uniform manifold approximation and projection (UMAP) plot on (C) all mice and (D) separated by age and sex, with cells color-coded based on SNN-clustering. Clusters 5 and 6 were enriched in MOTS-c-programmed BMDM populations. (E) Box plot of relative cluster cell proportion ratios between MOTS-c-programmed vs. control macrophages across clusters. Note that clusters 5 and 6 are consistently found in higher proportion in MOTS-c treated samples compared to their corresponding control condition. (F) Dotplot of select genes enriched in clusters 5 and 6 (see also Figures S10-S12 and Table S3). (G) Heatmap of the top 10 differential gene markers of each of the 8 clusters in BMDMs induced in the presence/absence of MOTS-c (FDR < 5%).
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Author contributions:

B.A.B. and C.L. conceived the experiments. M.C.R., J.S.K., M.I., S.W.J., C.Y.P., R.W.L., C.R.B., J.M.S., K.T., E.K., R.J.L., I.C., B.A.B., and C.L. performed experiments and analyzed the data. C.L. wrote the manuscript with input from authors. All authors approved the manuscript.

Competing interests:

C.L. is a consultant and shareholder of CohBar, Inc. All other authors declare no competing interests.
Methods

Bacterial Strains

*E. coli* [BL21 (DE3); NEB], MRSA (methicillin-resistant *S. aureus*) (ATCC 33592), *S. typhimurium* (ATCC 14028), and *P. aeruginosa* (ATCC 27853) were used. *E. coli* cells were transformed to conditionally overexpress MOTS-c in pSF-T7/LacO (Sigma, OGS500) using IPTG. Bacteria were routinely maintained in LB medium (liquid and agar). A modified M9 minimal medium, composed of 0.1x of M9 Minimal Salts without NaCl (BD Difco), 1 mM MgSO$_4$, 1% glucose, and 2.5 g/L peptone was used in select studies.

Phase-Contrast Light Microscopy

Bacteria were grown overnight in LB broth, from which a 1:100 inoculation was made in LB broth and grown to mid-log phase (~0.6 OD$_{600}$), measured using a SpectraMax M3 Spectrophotometer (Molecular Devices), in a 37°C shaker at 225 rpm. 1 mL of mid-log *E. coli* (BL21) and 3 mL of mid-log MRSA were collected and treated in modified M9 minimal medium to achieve macroscopic bacterial aggregates. Approximately 3 µL of aggregates were transferred to Superfrost Plus Micro Slides (VWR) with platinum-grade cover slips and imaged with phase-contrast mode using a EVOS FL Cell Imaging System.

Scanning Electron Microscopy

Samples were fixed overnight at 4 °C in 3% glutaraldehyde. Fixed cells were placed onto 0.2-µm Nuclepore Track-Etch Membrane filters (Whatman) and allowed to air dry for fifteen minutes prior to ethanol dehydration. The dehydration series progressed from an initial wash concentration of 30% ethanol with 30-minute stepwise increments to a final wash of 100% ethanol prior to critical point drying (Autosamdri-815, Toursimis). Samples were then sputter coated (Cressington) with approximately 3 nm of Pd. Electron micrographs were obtained with a JEOL-7001 FEG Scanning Electron Microscope.

Confocal imaging

Cellular images were obtained using a Zeiss LSM700 confocal microscope system (Germany). THP-1 monocytes were treated with synthetic MOTS-c peptide tagged with a FITC fluorophore (1 µM) for 30 min. After washing 3 times with PBS, cells were fixed using 4% paraformaldehyde and permeabilized in 0.2% TritonX100. Fixed cells were incubated with DAPI (Sigma) for 30 min and washed an additional three times with 0.1% PBST. The cells were spread
onto glass coverslips and attached to MicroSlides (cat#48311-703, VWR) using ProLong Gold antifade mountant (cat#P36934, ThermoFisher).

**Bacterial growth measurements**

Overnight bacterial cultures were diluted 1:1000 and grown in modified M9 minimal medium (0.1x M9, 5% glucose, 1mM MgSO4, 1.0 g/L peptone) with MOTS-c (or vehicle) in a 37°C shaker at 225 rpm. Bacterial density was measured at OD600 every hour in two-sided polystyrene cuvettes (VWR) using a SpectraMax M3 Spectrophotometer (Molecular Device). For plate assays, modified M9 minimal medium agar (1.5%) plates (35 mm Petri dishes) were made.

**ATP Assay**

BacTiter-Glo Microbial Cell Viability Kit (Promega) was used to assess bacterial cell viability and metabolism by measuring luminescence correlated with amount of intracellular ATP. Bacteria were grown overnight in LB broth, from which a 1:100 inoculation was grown in LB broth to reach mid-log phase (~0.6 OD600) in a 37°C shaker at 225 rpm. 1 mL of log-phase bacteria was collected, resuspended in modified M9 minimal medium, treated with MOTS-c (100 µM), then subject to luciferase reaction per the manufacturer’s instructions. Luminescence was measured using SpectraMax M3 Spectrophotometer (Molecular Device) and values reported as RLU (relative light units).

**SYTOX Green Assay**

SYTOX® Green Nucleic Acid Stain 5 mM in DMSO (ThermoFisher) was used to assess bacterial membrane integrity. SYTOX® Green does not cross intact membranes, but easily penetrates compromised membranes and stains nucleic acids52. Fluorescence readings (bottom-read, excitation/emission at 504/523 nm) were recorded at various time points with one measurement taken before treatment (blank).

**Western Blot**

Whole cell and nuclear compartment were lysed with 8 M Urea buffer containing a protease/phosphatase inhibitor cocktail (Thermo Fisher Scientific) and were sonicated for 15 seconds at 60% amplitude. The lysates were separated by 8-16% pre-cast SDS-PAGE gels (Bio-Rad) and transferred onto PVDF membranes. The membranes were blocked with 5% bovine serum albumin (BSA) in tris-buffered saline with 0.05% Tween-20 and probed with primary antibody at 4°C overnight. Proteins of interest were detected with anti-rabbit IgG HRP linked
antibody (Cell Signaling) and developed by Clarity Western ECL substrates (Bio-Rad). The membranes were imaged by the ChemiDoc XRS+ system (Bio-Rad).

**Human cell culture**

THP-1 cells (RRID:CVCL_0006) were routinely cultured at a range of cell density of 2\times10^5 cells/ml to 8\times10^5 cells/ml in RPMI 1640 (Corning) supplemented with 10% heat-inactivated fetal bovine serum (Omega Scientific) and 0.05 mM 2-Mercaptoethanol at 37°C with 5% CO₂. Macrophages were generated by culturing THP-1 (6\times10^5/ml) cells with 15 nM phorbol 12-myristate 13-acetate (PMA, Sigma-Aldrich) in the media. Human primary monocytes were isolated from leukocyte cones of healthy blood donors, obtained from USC/CHLA. Mononuclear cells were separated by centrifugation at 400\times g for 20 minutes with underlying Histopaque-1077. The opaque interface between plasma and the Histopaque-1077 was collected for further purification by Red Blood Cell lysis buffer (Miltenyi). Monocytes were isolated by negative selection using the StraightFrom LRSC CD14 MicroBead Kit (cat# 130-117-026, Miltenyi). Purified human monocytes were seeded at a density of 1\times10^6 cells/ml in RPMI 1640 supplemented with 10% heat inactivated fetal bovine serum and 100 ng/ml human M-CSF (Miltenyi).

**Human cell treatment**

THP-1 cells were seeded at a density of 4\times10^5 cells/ml and reached to 6\times10^5 cells/ml prior to treatment. PMA (Sigma-Aldrich) was reconstituted in DMSO and used at 15 nM. Lipopolysaccharides from Escherichia coli O111:B4 (List Biology Laboratories, Inc.) were reconstituted in sterile water and used at 100 ng/ml. Human interferon-gamma (Sigma) was reconstituted in sterile water and used at 20 ng/ml. Synthetic MOTS-c peptide (GenScript) was reconstituted in sterile water and used at 10 µM.

**Nuclear fractionation**

Nuclear fraction was isolated as previously described \(^{21,151}\). Cells were harvested by centrifugation. The pellet was resuspended in hypotonic fractionation buffer (10 mM Tris, pH 7.5, 10 mM NaCl, 3 mM MgCl₂ 0.3% NP-40, 10% glycerol, and EDTA-free protease inhibitor) and incubated on ice for 30 minutes and passed through a 31-gauge needle 5 times and centrifuged. The nuclear pellet was washed with hypertonic fractionation buffer 3 times and resuspend in 8 M urea lysis buffer (1 M Tris, pH = 8).
**Gentamicin Protection Assay**

THP-1 monocytes were seeded on a 6-well plate at concentration of 1.5x10^6 cells/well and differentiated by PMA (15 nM) for 96 hours. For MOTS-c treatment, monocytes were first primed with MOTS-c (10 μM) for 2 hours, then treated only once with a mixture of PMA (15 nM) and MOTS-c (10 μM). *E. coli* (BL21) were added to differentiated THP-1 cells (MOI 10) and co-cultured for 1-2.5 hours, at which time gentamicin (50 µg/ml) (Fisher Scientific) was added. 30 minutes later, cells were washed with PBS and collected (total of 1.5 or 3 hours). Cells were lysed with 1% Triton X-100 and spread on LB agar plate. Bacterial colonies were counted after overnight incubation at 37°C.

**ELISA**

THP-1 monocytes (n=6) were cultured in six-well plates at a density of 5 x 10^6 cells/mL in 2 mL of RPMI 1640 (cat#45000-396, VWR, USA) supplemented with 10% fetal bovine serum (Omega Scientific) and incubated at 37°C in humidified air with 5% CO₂. THP-1 cells were primed with MOTS-c peptide (10 μM) for 2 hours, then treated with a combination of PMA (15 nM) + MOTS-c (10 μM). 96 hours post-differentiation, cells were treated with LPS (100 ng/mL) or vehicle for an additional 20 hours and supernatant were collected for cytokine analysis. The production levels of cytokines including IL-1β (cat#BMS224-2), IL-1Ra (cat#BMS2080), and TNF-α (cat#BMS223-4) were measured in duplicate using human ELISA kits following the manufacturer’s instructions (ThermoFisher) and SpectraMax M3 (Molecular devices).

**Real-time qPCR**

Total RNA was extracted and purified using the Direct-zol RNA MiniPrep kit (Zymo Research) following manufacturer’s instructions. The total RNA (1.2 μg) was used to synthesize single-stranded cDNA using iScript cDNA synthesis Kit (Bio-Rad) and T100 Thermal Cycler (Bio-Rad) according to the manufacturer’s instructions. The relative mRNA levels of CCL2, CCL3, CCL5, CXCL9, CXCL10, CXCL11, IL-6, IL-10, IL-12A, and IL-12B were determined by the real-time quantitative PCR analysis using the SYBR Green Supermix (#1725275, Bio-Rad) and CFX Connect Real-time PCR Detection System (Bio-Rad). All reactions were performed in total of 20 μL reaction volume containing 2 μL of diluted cDNA, 10 μL of SYBR Supermix, 1 μL of 10 μM forward primer, 1 μL of 10 μM reverse primer, and 6 μL of autoclaved distilled water. All samples were analyzed in duplicate with an endogenous control gene (β-Actin) being analyzed at the same time. The relative expression of each gene was calculated from 2^ΔΔCT.
Primer sequences used in real-time RT-qPCR

<table>
<thead>
<tr>
<th>Gene</th>
<th>Forward primer</th>
<th>Reverse primer</th>
</tr>
</thead>
<tbody>
<tr>
<td>IL-12A</td>
<td>GATGGCCCTGTGCCTTAGTA</td>
<td>TCAAGGGAGGATTTTGTGG</td>
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<tr>
<td>IL-12B</td>
<td>GGACATCATCAAAACCTGACC</td>
<td>AGGGAGAAGTAGGAGATGTGG</td>
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<tr>
<td>IL-10</td>
<td>GTGATGCCCCAAGCTGAGA</td>
<td>CACGCGCTTGTGCTTTTGG</td>
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<tr>
<td>IL-6</td>
<td>TGCCTCCGTAGTGTTCCTTCT</td>
<td>GCCTCGACATCTCCAGTCC</td>
</tr>
<tr>
<td>CCL2</td>
<td>ATCAATGCCCCAGTCACC</td>
<td>AGTCTTCGGAGTGTGGG</td>
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<td>CCL3</td>
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<td>TACCATTAGGTGTCCGC</td>
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<td>CXCL9</td>
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<tr>
<td>CXCL11</td>
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</tbody>
</table>

Metabolic flux measurements

Real-time analysis of oxygen consumption rates (OCR) were measured in THP-1 cells using XF24/96 Extracellular Flux Analyzer (Seahorse Bioscience). Cells were seeded in Seahorse XF96 cell culture microplate (#101085-004, Seahorse Bioscience) at a density of 5 x 10^6 cells/mL in 100 μL of RPMI 1640 (#45000-396, VWR) supplemented with 10% fetal bovine serum (Omega Scientific). The cells were treated without or with 10 μM MOTS-c for 2 hours, followed by treatment with PMA for 96 hours. LPS (100 ng/mL) was added either during OCR measurements (dispensed by XF96) or added for 16 hours prior to OCR measurements and medium was replaced with XF assay buffer supplemented with 1 mM pyruvate and 12 mM D-glucose. ATP turnover, maximum respiratory capacity, and non-mitochondrial respiration were estimated by sequential addition of oligomycin (0.9 μM), carbonyl cyanide 4-[trifluoromethoxy]phenylhydrazone (FCCP, 1.0 μM), rotenone (0.5 μM), and antimycin A (0.5 μM). All readings were normalized to relative protein concentration.

Bulk RNA-seq library preparation

1 μg of total RNA was subjected to rRNA depletion using the NEBNext rRNA Depletion Kit (New England Biolabs), according to the manufacturer’s protocol. Strand specific RNA-seq libraries were then constructed using the SMARTer Stranded RNA-Seq Kit (Clontech #634839),...
according to the manufacturer’s protocol. Based on rRNA-depleted input amount, 13-15 cycles of amplification were performed to generate final RNA-seq libraries. Pooled libraries were sent for paired-end sequencing on the Illumina HiSeq-Xten platform at the Novogene Corporation (USA). The raw sequencing data has been deposited to the NCBI Sequence Read Archive (accession number PRJNA623667). The resulting data was analyzed with a standard RNA-seq data analysis pipeline (described below).

**Bulk RNA-seq analysis pipeline**

To avoid the mapping issues due to overlapping sequence segments in paired-end reads, reads were hard trimmed to 75bp using Fastx toolkit v0.0.13. Reads were then further quality-trimmed using Trimgalore 0.4.4 to retain high-quality bases with Phred score > 20. All reads were also trimmed by 6 bp from their 5’ end to avoid poor qualities or random-hexamer driven sequence biases. cDNA sequences of protein coding and IncRNA genes were obtained through ENSEMBL Biomart for the GRCh38 build of the human genome (release v96). Trimmed reads were mapped to this reference using kallisto v0.43.0 and the –fr-stranded option. All bulk RNA-seq analysis were performed in the R statistical software Version 3.4.1 ([https://cran.r-project.org/](https://cran.r-project.org/)). Read counts were imported into R, and summarized at the gene level to estimate gene expression levels. We estimated differential gene expression between control, PMA and PMA+MOTS-c treated THP-1 RNA-seq samples using the ‘DESeq2’ R package (DESeq2 1.16.1). The heatmap of expression across samples for significant genes ([Figure 5D](#)) was plotted using the R package ‘pheatmap’ 1.0.10.

**Functional enrichment analysis**

To perform functional enrichment analysis, we used the Gene Set Enrichment Analysis (GSEA) paradigm through R packages ‘phenoTest’ 1.24.0 and ‘qusage’ 2.10.0. Gene Ontology (GO) gene sets were obtained from the Molecular Signature Database, C5 collection (c5.all.v6.2.symbols.gmt). A False Discovery Rate (FDR) threshold of 0.05 was considered statistically significant. The -log10(FDR) value for GSEA enrichment is reported in Figure 3F as a barplot for selected terms, and FDR values of 0 were replaced by a small value of 10^{-30} to enable plotting on a reasonable scale. The full list of enriched GO terms at FDR 0.05 is reported in Table S2.
STRING analysis

The list of significantly regulated genes in the PMA vs. PMA+MOTS-c conditions at FDR < 0.05 was used to infer potential disruption to protein interaction networks using the STRING (Search Tool for the Retrieval of Interacting Genes/Proteins) tool database version 11.0.

Mouse husbandry

All animals were treated and housed in accordance with the Guide for Care and Use of Laboratory Animals. All experimental procedures were approved by the University of Southern California's Institutional Animal Care and Use Committee (IACUC) and are in accordance with institutional and national guidelines. For murine peritonitis experiments, male C57BL/6J mice were obtained from Jackson Laboratory and experiments performed at 14-16 weeks of age. For single-cell RNA-seq analyses, male and female C57BL/6JNia mice (4 and 20 month old animals) were obtained from the National Institute on Aging (NIA) colony at Charles Rivers. Animals were acclimated at the SPF animal facility at USC for 2-4 weeks before any processing, and were euthanized between 8-11am to minimize circadian effects. In all cases, animals were euthanized using a “snaking order” across all groups to minimize batch-processing confounds. All animals were euthanized by CO₂ asphyxiation followed by cervical dislocation.

Murine model of peritonitis

MRSA (methicillin-resistant *S. aureus*; ATCC 33592) were grown overnight in LB broth, from which a 1:100 inoculation was made in LB broth and grown to mid-log phase (~0.6 OD₆₀₀), measured using a SpectraMax M3 Spectrophotometer (Molecular Devices), in a 37°C shaker at 225 rpm. 4x10⁸ CFU of MRSA was washed and resuspended in 100 µM MOTS-c or water immediately prior to intraperitoneal injection with a 28G syringe. MRSA was serial diluted, plated on LB agar, and colonies counted after overnight incubation to confirm CFU inoculated. Mice were monitored for 72 hours and weighed daily. Moribund animals were euthanized by CO₂ asphyxiation followed by cervical dislocation according to IACUC approved experimental guidelines. To heat-kill MRSA, MRSA resuspended in phosphate-buffered saline (PBS) was heated in a 70°C water bath for 20 minutes.
Derivation of bone marrow-derived macrophages (BMDMs) and MOTS-c treatment

We isolated BMDMs as previously described. Briefly, the long bones of each mouse were harvested and kept on ice in D-PBS (Corning) supplemented with 1% Penicillin/Streptomycin (Corning) until further processing. Muscle tissue was removed from the bones, and the bone marrow from cleaned bones was collected into clean tubes. Red blood cells from the marrow were removed using Red Blood Cell Lysis buffer (Miltenyi Biotech #130-094-183), according to the manufacturer's instructions, albeit with no vortexing step and incubation for only 2 minutes. The suspension was filtered on 70 μm mesh filters to retain only single cells. Cells were plated in macrophage growth medium [DMEM/F12 (Corning), 10% FBS (Sigma), 1% Penicillin/Streptomycin (Corning), 2 ng/mL recombinant M-CSF (Miltenyi Biotech), and 10% L929-conditioned medium as an additional source of M-CSF]. For cells undergoing MOTS-c treatment, this initial culture medium was supplemented with 10 μM MOTS-c peptide. After 3 days, adherent cells were rinsed with D-PBS, and fresh macrophage growth media was added. Cells were collected after 7 days in culture, when BMDMs have completed differentiation. We differentiated BMDMs from 5 animals from each group (young females, young males, old females and old males, control and MOTS-c treated; 8 groups total).

BMDM single-cell RNA-seq library preparation

Single-cell RNAseq libraries were prepared using Single Cell 3’ v2 Reagent Kits, according to the manufacturer's instructions (10xGenomics). For single-cell RNA-seq profiling, BMDMs were detached using ice-cold 10 mM EDTA, counted using a COUNTESS cell counter (Thermo Fisher Scientific), and samples from each sample group were pooled in an equicellular mix (8 mixes, 1 mix per biological condition). Using the 10xGenomics Single Cell 3’ v2 manufacturer's instructions (10xGenomics), we loaded the microfluidics device with a targeted capture in each sample of 3,000 cells. The 8 samples were run in parallel on the same microfluidics chip and processed on a Chromium Controller instrument (10xGenomics), to generate single-cell Gel bead-in-EMulsions (GEMs). GEM-RT was performed in a C1000 Touch Thermal Cycler with a deep well module (Biorad). The cDNA was amplified and cleaned up with SPRIselect Reagent Kit (Beckman Coulter Genomics). Single-indexed sequencing libraries were constructed using Chromium Single-Cell 3’ Library Kit. Library quality and quantification prior to pooling were assessed using a D1000 screentape device on the Tapestation apparatus (Agilent). All barcoded libraries were combined in a single pool for sequencing, and sent for sequencing on 3 lanes of Hiseq-X-Ten at Novogene.
Corporation as paired end 150bp reads. The final average sequencing depth per cell was ~70,000 reads per cell.

**BMDM single-cell RNA-seq data analysis**

Reads were hard trimmed to yield the lengths expected by the CellRanger pipeline (Read1: 26bp, Read 2: 98bp) using the fastx_trimmer tool from the FASTX Toolkit v0.0.13 [http://hannonlab.cshl.edu/fastx_toolkit/]. The raw sequencing data has been deposited to the NCBI Sequence Read Archive (accession number PRJNA769064). Trimmed reads were then processed using CellRanger software version 3.0.2 and the mm10 mouse genome reference for mapping, cell identification and UMI processing (10X Genomics). Analyses for single-cell RNA-seq were performed using R version 3.6.3 (single-cell level clustering and marker identification) or 4.1.2 (pseudobulk analysis).

Since sequencing on new generation Illumina patterned flow cells can lead to the emergence of non-physiological chimeric reads, phantom molecules were identified and removed from the cellranger h5 output using the PhantomPurge 1.0.0 R package. After purging, data was converted to the ‘Seurat’ format for single cell analysis for analysis with Seurat 3.2.2 package. Genes detected in at least 50 cells, and cells with greater than 1000 genes and less than 10% of mitochondrial genes were selected for downstream analysis, yielding 15,415 cells and 11,622 genes passing quality control filters. The impact of number of genes detected per cells, percentage mitochondrial read and cell cycle phase were regressed out as recommended by the Seurat package. We next leverage the DoubletFinder 2.0.3 package to identify and filter out likely cell doublets, assuming a 2.3% doublet formation rate based on 10xGenomics estimates when capturing 3,000 cells per samples. After this, 15,060 cells were identified as singlets by DoubletFinder and used for downstream analyses. Data was normalized using the SCtransform framework. We then ran Principal Component Analysis (PCA), dimensionality reduction using UMAP using 30 principal components, and clustering with a resolution set to 0.2, yielding a total of 8 clusters. Cluster markers were identified using the “FindAllMarkers” function with a minimum 25% of cells and a log2 fold change threshold of 0.25 using the Wilcoxon-test, and a significance threshold of FDR < 0.05.

To analyze the scRNA-seq dataset with a pseudobulk approach, unnormalized counts were aggregated from cells within the same group (8 groups across age, sex, and MOTS-c treatment). Using R version 4.1.2 and the DESeq2 1.34.0 R package, differential transcriptomic
profiles between the groups was estimated and visualized with multidimensional scaling (MDS). To estimate the pairwise distance between female and male pseudobulk samples, we used a distance metric based on Spearman Rank Correlation Rho (1-Rho). The distance of each female sample to both male samples was calculated in the control and MOTS-c treated conditions. To determine whether the distance between female and male samples was reduced upon MOTS-c treatment (as hypothesized based on the MDS analysis), we then used a paired one-sided Wilcoxon Rank sum test to compare the distances in control vs. MOTS-c treated conditions.

**BMDM single cell Gene Ontology cluster marker data analysis**

To analyze which functional categories were enriched in association to BMDMs in clusters 5 and 6, whose frequency increased upon MOTS-c treatment, we used overrepresentation analysis with clusterProfiler 3.14.3 and annotation package org.Mm.eg.db 3.10.0. Enrichment was computed against the background of all expressed genes detected in the single cell dataset.

**Code availability**

The analytical code for the RNA-seq dataset is available on the Benayoun lab github (https://github.com/BenayounLaboratory/MOTSc_Macrophage_Immunity). R code was run using R version 3.4.1, 3.6.3 or 4.1.2 as indicated in relevant sections.
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Figure 2

A-C

Control
MOTS-c
MRSA

A

Survival (%)

0 12 24 36 48 60 72

P=0.026

0 24 48 72

Hours

B

Weight (%)

80 85 90 95 100

P=0.0079

0 24 48 72

Hours

C

CFU Inoculated (x10^7)

Control
MOTS-c

P=0.0079

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Figure 3

A

M-CSF: 0 (hrs) 20 4 40 (hrs)
IB: MOTS-c
IB: β-actin

Primary Human Monocytes:

B

PMA: 0 8 24 48 (hrs)
IB: MOTS-c
IB: β-actin

THP-1 Monocytes

C

LPS+IFNγ: 0 4 8 24 (hrs)
IB: MOTS-c
IB: β-actin

D

LPS: 0 4 8 24 (hrs)
IB: MOTS-c
IB: β-actin

E

IFNγ: 0 4 8 24 (hrs)
IB: MOTS-c
IB: β-actin

MOTS-c/β-actin (a.u.)

0 4 20 40 (hrs)

0 24 8 4 (hrs)

0 24 8 4 (hrs)

0 24 8 4 (hrs)

0 48 24 8 (hrs)