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Aiolos promotes CXCR3 expression on T_H1 cells via positive regulation of IFNγ/STAT1 signaling

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1	Title: Aiolos promotes CXCR3 expression on T_H1 cells via positive regulation of
2	IFNy/STAT1 signaling
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26 Conflict-of-interest statement

27 The authors have declared that no conflict of interest exists.

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29 Abstract: CD4⁺ T helper 1 (T_H1) cells coordinate adaptive immune responses to intracellular 30 pathogens, including viruses. Key to this function is the ability of $T_{\rm H}1$ cells to migrate within 31 secondary lymphoid tissues, as well as to sites of inflammation, which relies on signals received through the chemokine receptor CXCR3. CXCR3 expression is driven by the T_H1 lineage-defining 32 transcription factor T-bet, and the cytokine-responsive Signal Transducer and Activator of 33 34 Transcription (STAT) family members STAT1 and STAT4. Here, we identify the lkaros zinc finger (IkZF) transcription factor Aiolos (Ikzf3) as an additional positive regulator of CXCR3 both in vitro 35 36 and in vivo using a murine model of influenza virus infection. Mechanistically, we find that Aiolos-37 deficient CD4⁺ T cells exhibit decreased expression of key components of the IFNy/STAT1 38 signaling pathway, including JAK2 and STAT1. Consequently, Aiolos deficiency results in decreased levels of STAT1 tyrosine phosphorylation and reduced STAT1 enrichment at the Cxcr3 39 promoter. We further find that Aiolos and STAT1 form a positive feedback loop via reciprocal 40 regulation of each other downstream of IFNy signaling. Collectively, our study demonstrates that 41 42 Aiolos promotes CXCR3 expression on T_H1 cells by propagating the IFNy/STAT1 cytokine signaling pathway. 43

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52 Introduction

53 During adaptive immune responses to infection, naïve CD4⁺ T cells differentiate into T "helper" subsets with distinct effector functions. T helper 1 (T_{H} 1) cells represent one such subset that 54 produces interferon gamma (IFNy) and directs immune responses against intracellular 55 56 pathogens. Differentiation of effector subsets is initiated when antigen-presenting cells (APCs) 57 deliver cognate antigen to the T cell receptor (TCR) expressed on the surface of a naïve CD4⁺ T cell. Additional signals received in the form of co-stimulatory molecules and environmental 58 cytokines further propagate T cell activation and differentiation into specific subsets (1-5). As part 59 60 of this process, signaling through cytokine receptors leads to activation of Janus kinases (JAKs) and ensuing tyrosine phosphorylation of STAT factors, which then dimerize, translocate to the 61 62 nucleus, and bind to target genes (6-9). For example, IFNy signaling via STAT1 and IL-12 signaling via STAT4 both stimulate the expression of T-bet (Tbx21), the lineage-defining 63 64 transcription factor for the $T_H 1$ gene program (10-12). In turn, T-bet promotes IFNy production via direct transcriptional activation of the *lfng* gene, thus, creating a positive feedback loop that drives 65 T_{H1} differentiation (13-15). 66

More recently, Ikaros Zinc Finger (IkZF) transcription factors have been implicated in the 67 68 regulation of CD4⁺ T cell programming events (16, 17). IkZF factors contain conserved N-terminal zinc finger (ZF) domains that mediate DNA-binding specificity as well as C-terminal protein-69 interaction domains that enable IkZF dimerization and recruitment of co-regulators, such as 70 71 chromatin remodeling complexes (16-22). The importance of IkZF factors to immune cell function 72 is underscored by studies in humans that have described missense mutations in the IkZF family 73 member Aiolos (Ikzf3) that result in immunodeficiency. These changes have been associated with 74 abnormalities in B and T cell differentiation, increased susceptibility to infectious diseases, and elevated risk for certain types of hematological malignancies (23-26). One study specifically 75 76 identified a heterozygous Aiolos mutation associated with impaired T helper cell polarization, 77 which led to reduced numbers of T follicular helper (T_{FH}) and T_{H1} cells in affected patients (25).

78 IkZF factors have also emerged as key regulators of cytokine signaling pathways (18). 79 Previous work from our lab identified a transcription factor complex comprised of Aiolos and STAT3 that promoted T_{FH} cell programming (18). Similarly, we established that a second 80 81 IkZF/STAT factor complex comprised of Eos (*Ikzf4*) and STAT5 drives T_{H2} differentiation by 82 inducing expression of IL-4 and IL-2 cytokine receptors (27). Further, we demonstrated that Aiolos modulates IL-2 responsiveness via repression of the IL-2R α (CD25) and IL-2R β (CD122) 83 subunits, both promoting T_{FH} differentiation and suppressing cytotoxic programming of CD4⁺ T 84 cells (28). Beyond our findings, others have found that Aiolos regulates cytokine production, 85 including direct silencing of the *II2* locus in $T_H 17$ cells (29). These collective findings suggest that 86 87 a complex interplay exists between IkZF factors, cytokine signaling pathways, and T helper cell 88 programming events, much of which remains enigmatic.

Like cytokines, chemokines signal through specific receptors, and serve as integral 89 mediators of both CD4⁺ T cell differentiation and migration (30-32). The chemokine receptor 90 CXCR3 is a G protein-coupled receptor that is highly expressed on the surface of T_{H1} cells (33, 91 34). CXCR3 responds to three interferon-inducible ligands (CXCL9, CXCL10, and CXCL11) and 92 93 directs T_H1 cells to sites of inflammation (33, 35, 36). Although T_H1 cell responses are beneficial during infection, their activities are typically tightly controlled to prevent destruction of healthy 94 95 tissue. To this end, aberrant activities of T_H1 cells have been implicated in autoimmunity, and 96 therapeutically targeting CXCR3 appears to have disease-specific advantages (35, 37-43). As 97 such, obtaining a better understanding of the transcriptional mechanisms regulating CXCR3 98 expression may provide additional insight for the development of novel therapeutics.

99 Here, we identify Aiolos as a positive regulator of CXCR3 expression in both in vitro-100 generated T_H1 cells and those that arise in response to murine influenza virus infection. 101 Mechanistically, we find that Aiolos-deficient CD4⁺ T cells have reduced expression of 102 components of the IFNγ/STAT1 signaling pathway, which results in decreased STAT1 activation 103 and enrichment at the *Cxcr3* promoter. We further find that *Stat1* is a direct Aiolos target gene,

104	with Aiolos both modulating chromatin accessibility at, and driving activity of, the Stat1 promoter.
105	Moreover, we demonstrate that Aiolos expression itself is dependent upon IFN γ signaling and
106	that STAT1 directly binds the <i>lkzf3</i> promoter. Collectively, our findings reveal that Aiolos promotes
107	CXCR3 expression on $T_{\text{H}}1$ cells by propagating IFNγ/STAT1 signaling via a positive feedback
108	loop with STAT1.
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130 **Results**

131 CXCR3 expression is reduced on Aiolos-deficient CD4⁺ T_H1 cells.

We previously reported that Aiolos functions as a repressor of CD4⁺ T cell cytotoxic programming 132 via suppression of IL-2/STAT5 signaling (28). In line with these findings, RNA-seg analysis of 133 134 wild-type (WT) versus Aiolos-deficient ($Ikzf3^{-/}$) T_H1 cells revealed increased expression of many 135 STAT5 target genes associated with cytotoxic function, including Gzmb, Prf1, Ifng, and Prdm1 (Figure 1A). In contrast, expression of Cxcr3 (encoding CXCR3), a chemokine receptor that 136 guides migration of both T_H1 cells and CD4⁺ cytotoxic T lymphocytes (CD4-CTLs), was reduced 137 138 in the absence of Aiolos (Figure 1A). Consistent with the RNA-seq data, transcript and flow cytometric analyses of in vitro-generated $lkzf3^{-/-}T_H1$ cells revealed significant decreases in both 139 CXCR3 transcript and cell surface expression compared to WT after 3 days of differentiation 140 141 (Figure 1, B-D). In accordance with previous reports describing the inhibition of CXCR3 142 expression with persistent TCR stimulation, the reduction in cell surface expression of CXCR3 on *lkzf3^{-/-}* T_H1 cells was further enhanced after cells were removed from stimulation with α -CD3 and 143 144 α -CD28 antibodies and cultured for an additional 48 hours (Figure 1, B, E, F) (44). These findings suggested that CXCR3 expression may be regulated by an Aiolos-dependent mechanism. 145

146 We next examined the impact of Aiolos deficiency on CXCR3 expression in vivo using a murine model of influenza A virus (IAV) infection (45). WT and *lkzf3^{-/-}* mice were intranasally 147 infected with a sublethal dose of IAV strain A/PR/8/34 (H1N1, termed "PR8"), and nucleoprotein 148 (NP)-specific CD4⁺ T cells of the draining, mediastinal lymph node (mLN) and lungs were 149 assessed at 8 days post-infection (Figure 2A). There was no significant difference in the numbers 150 of NP-specific CD4⁺ T cells in the mLN between WT and *lkzf3^{-/-}* mice (Supplemental Figure 1A). 151 Further analyses of the NP-specific population revealed a significant decrease in CXCR3 surface 152 expression in the absence of Aiolos (Figure 2, B and C). In contrast to the mLN, *lkzf3^{-/-}* mice had 153 154 a significant reduction in NP-specific CD4⁺ T cells in the lungs compared to WT (Supplemental

155 Figure 1B). These findings are in agreement with previous reports identifying CXCR3 as an 156 essential chemokine receptor for antigen-specific effector T cell recruitment to the lungs (46, 47). 157 The numbers of bulk CD4⁺ T cells in the mLN were quantified and again revealed no significant difference between WT and *lkzf3^{-/-}* mice (Supplemental Figure 1C). However, in the lunas, we 158 159 observed a significant reduction in the number of bulk CD4⁺ T cells in *lkzf3^{-/-}* mice compared to WT, though not to the extent observed with NP-specific cells (Supplemental Figure 1D). 160 Collectively, these data suggest that migration of CD4⁺ T cells is disrupted in Aiolos-deficient mice 161 162 during pulmonary infection.

163 T-bet is a known positive regulator of CXCR3 expression (48-51). Thus, we next examined 164 T-bet expression to determine whether differences in this transcriptional regulator may explain 165 the decrease in CXCR3 expression. However, we observed no significant difference in T-bet 166 expression between NP-specific WT and $lkzf3^{-/-}$ cells, suggesting that Aiolos-dependent 167 regulation of CXCR3 expression may occur through a T-bet-independent mechanism 168 (Supplemental Figure 2A).

Finally, we examined CXCR3 surface expression on bulk CD4⁺ naïve, central memory, 169 170 and effector T cell populations from the spleen to determine whether the decrease in CXCR3 expression was limited to distinct CD4⁺ T cell subsets. As expected, naïve CD4⁺ T cells from both 171 WT and *lkzf3^{-/-}* mice did not express CXCR3 (Supplemental Figure 2B). However, both central 172 memory and effector CD4⁺ T cells from $lkzf3^{-/-}$ mice displayed significantly reduced CXCR3 173 expression compared to their WT counterparts, with a greater reduction present on effector CD4⁺ 174 T cells (Supplemental Figure 2, C and D). Overall, our findings in NP-specific and bulk CD4⁺ T 175 cells suggest that Aiolos regulates CXCR3 expression in vivo. 176

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178 CXCR3 expression is reduced on Aiolos-deficient CD4⁺ T cells in a cell-intrinsic manner.

To assess whether the impact of Aiolos on CXCR3 expression was CD4⁺ T cell-intrinsic, we crossed WT and $lkzf3^{-/-}$ mice onto the OT-II background, which expresses a transgenic TCR

specific for the ovalbumin 323-339 peptide (OVA). We then adoptively transferred naïve CD45.2+ 181 CD4⁺ T cells from either WT-OT-II or *lkzf3^{-/-}*-OT-II mice into WT CD45.1⁺ recipients. Recipient 182 mice were subsequently infected with OVA323-339-expressing PR8 (PR8-OVA) 24 hours post-183 184 transfer and antigen-specific CD45.2⁺ donor cells from the mLN were analyzed via flow cytometry 185 at 8 days post-infection (Figure 3A). Consistent with our findings in Aiolos-deficient mice, we observed a significant decrease in CXCR3 expression on donor CD45.2⁺ *lkzf3^{-/-}* cells compared 186 to WT (Figure 3, B and C). We also noted a slight downregulation in T-bet expression in *Ikzf3^{-/-}* 187 compared to WT CD45.2⁺ donor cells (Supplemental Figure 3A). However, the fold reduction in 188 189 CXCR3 expression was greater than that of T-bet. Collectively, these findings demonstrate that the impact of Aiolos on CXCR3 expression occurs in a CD4⁺ T cell-intrinsic manner. 190

We next quantified the numbers of antigen-specific CD45.2⁺ donor cells in the mLN and 191 192 lungs of recipient mice. In the mLN, there was no significant difference between the number of WT-OT-II and *lkzf3^{-/-}*-OT-II cells, and only a slight but significant decrease in the frequency of 193 Aiolos-deficient cells (Supplemental Figure 3B). In contrast to our findings in germline knockout 194 195 animals, there was no significant difference in cell numbers or percentages between donor WT-OT-II and *lkzf3^{-/-}*OT-II cells in the lungs (Supplemental Figure 3C). These data indicate that the 196 197 migration of donor T cells to the lungs is not impaired when cells are intravenously injected into 198 recipient mice, despite the observed reduction in CXCR3 expression. Thus, while Aiolos regulates CXCR3 expression in CD4⁺ T cell-intrinsic manner, alterations in CXCR3 expression do not 199 200 ultimately have a cell-intrinsic effect on migration of adoptively transferred cells to the lungs.

To determine a potential explanation, we analyzed published RNA-seq data (GSE203065) from WT and *lkzf3^{-/-}* T_H1 cells. As with *Cxcr3*, various adhesion molecules and integrin subunits were downregulated in the absence of Aiolos. However, several other chemokine receptors known to promote T_H1 cell migration (i.e. *Ccr5*, *Cxcr6*) were upregulated in Aiolos-deficient cells relative to WT (Supplemental Figure 3D). Hence, the impact of Aiolos on migratory programming

appears to be multilayered with alterations to other migratory receptors potentially compensating
 for the loss of CXCR3 in the adoptive transfer setting.

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Aiolos deficiency alters expression of components of the IFNγ/STAT1 and IL-12/STAT4 pathways.

211 We next sought to identify the mechanism(s) by which Aiolos may regulate CXCR3 expression in T_H1 cells. In addition to T-bet, CXCR3 expression is regulated by STAT1 and STAT4 (44, 52-54). 212 213 Comparison of publicly available Chromatin Immunoprecipitation sequencing (ChIP-seq) data for 214 STAT1 (GSM994528), STAT4 (GSM550303), and T-bet (GSM836124) revealed enrichment of these factors at the promoter and 3' enhancer regions of the Cxcr3 locus (Figure 4A) (48-51, 55-215 58). Analysis of published RNA-seq data (GSE203065) from WT and *Ikzf3^{-/-}*T_H1 cells showed that 216 217 the expression of key components of the IFNy/STAT1 and IL-12/STAT4 signaling pathways was 218 altered in the absence of Aiolos. Specifically, Jak2 and Stat1 were downregulated in $lkzf3^{-1}$ compared to WT T_H1 cells (Figure 4B) (28). Notably, Jak2, which is shared between the IFNy and 219 IL-12 pathways, was the only Janus kinase that was significantly downregulated in $lkzf3^{-/-}T_{H1}$ 220 221 cells (findings available in Source Data). Further, genes encoding the IFNy and IL-12 cytokine 222 receptor subunits, Ifngr2 and II12rb1, also displayed slight decreases in the absence of Aiolos 223 (Figure 4B). Given that many of the altered genes encode proteins involved in IFNy/STAT1 and IL-12/STAT4 signaling, we hypothesized that Aiolos may regulate CXCR3 via impacts on these 224 225 pathways (Figure 4C).

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227 IFNγ/STAT1 signaling, but not IL-12/STAT4, is diminished in the absence of Aiolos.

We next assessed the impact(s) of Aiolos on IFN γ /STAT1 and IL-12/STAT4 signaling with WT and *lkzf3^{-/-}*CD4⁺ T cells cultured under T_H1 conditions. Previous work from our lab has established that Aiolos-deficient T_H1 cells have enhanced IFN γ production upon stimulation (28). To control for this, as well as any other cytokines produced by CD4⁺ T cells in culture, cells were removed 232 from stimulation on day 3 of differentiation and cultured for an additional 2 days with IL-12 before 233 being harvested for analysis (Figure 5A). Transcript analysis of IL-12-treated cells revealed significant decreases in Cxcr3, Stat1, Jak2, and Ifngr2 in Ikzf3^{-/-} relative to WT T_H1 cells. In 234 235 contrast, transcript levels for Stat4 and II12rb2 were significantly increased in Ikzf3^{-/-}T_H1 cells 236 compared to WT (Figure 5B and Supplemental Figure 4A). No significant difference in transcript 237 for Tbx21 was observed between groups (Figure 5B). Immunoblot analyses similarly revealed significant reductions in total JAK2 and STAT1 protein in the absence of Aiolos (Figure 5, C and 238 D). In contrast, tyrosine-phosphorylated STAT4 (pY-STAT4) and total STAT4 protein levels were 239 240 significantly elevated in the absence of Aiolos (Figure 5C), demonstrating that the STAT4 pathway is not functionally inhibited by Aiolos deficiency when cells are treated with IL-12. 241

To more directly assess the impact of Aiolos-deficiency on the IFNy/STAT1 signaling 242 243 pathway, we cultured WT and $lkzf3^{-/2}$ CD4⁺ T cells under T_H1 conditions for 3 days, and then added 244 IFNy, rather than IL-12, for an additional 48 hours in the absence of stimulation (Figure 6A). Again, *lkzf3^{-/-}* T_H1 cells had significantly reduced expression of *Cxcr3*, *Stat1*, *Jak2*, and *lfngr2* compared 245 246 to WT, whereas Stat4 and II12rb2 transcripts were consistently increased (Figure 6B and 247 Supplemental Figure 4B). In contrast to IL-12-treated cells, we also found that Tbx21 expression 248 was significantly decreased in Aiolos-deficient cells, suggesting that IFNy/STAT1 signaling is 249 unable to compensate for the lack of IL-12/STAT4-dependent activation of T-bet expression in the absence of Aiolos (Figure 6B) (54). 250

Flow cytometry analysis similarly revealed a significant decrease in CXCR3 surface expression on *lkzf3^{-/-}* cells compared to WT (Figure 6C). Immunoblot analyses showed significant reductions in total JAK2, tyrosine-phosphorylated STAT1 (pY-STAT1), and total STAT1 protein in the absence of Aiolos (Figure 6D). However, total STAT4 protein remained significantly elevated in *lkzf3^{-/-}* T_H1 cells (Supplemental Figure 4C). Thus, across IL-12 and IFNγ simulation conditions, only JAK2 and STAT1 correlated with the reduced CXCR3 expression observed in the absence of Aiolos.

258 We next examined publicly available STAT1 ChIP-seq data (GSM994528) to identify 259 STAT1-binding regulatory regions at the *Cxcr*3 locus that may be impacted by Aiolos-deficiency and performed ChIP analysis of IFNy-treated WT versus $lkzf3^{-/-}$ T_H1 cells (Figure 6E) (56). 260 Coincident with the loss of STAT1 expression, we observed a significant decrease in STAT1 261 262 enrichment at the Cxcr3 promoter and a trending decrease in STAT1 enrichment at the Cxcr3 3' 263 enhancer region (p = 0.0697) in the absence of Aiolos (Figure 6F). These collective data demonstrate that the IFNy/STAT1 pathway is compromised by Aiolos deficiency and that Aiolos 264 functions to positively regulate CXCR3 expression via STAT1. 265

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267 IFNγ/STAT1 signaling induces Aiolos expression.

268 We next wanted to examine how directly inhibiting IFNy signaling may impact the relationship 269 between Aiolos, STAT1, and CXCR3. We cultured WT naïve CD4⁺ T cells under T_H1 conditions 270 for 3 days with the addition of an IFNy-neutralizing antibody to inhibit autocrine IFNy signals (Figure 7A). Neutralizing IFNy led to significant reductions in Cxcr3, Stat1, and Jak2 expression 271 272 (Figure 7B) (44, 52). Flow cytometric and immunoblot analyses similarly revealed significant 273 reductions in CXCR3, total JAK2, total STAT1, and pY-STAT1 protein levels when IFNy was 274 neutralized (Figure 7, C and D). Notably, Aiolos expression was also significantly decreased with IFNy neutralization, suggesting a possible positive feedback loop between Aiolos and 275 IFNy/STAT1 (Figure 7E). 276

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Aiolos and STAT1 are enriched at the *Stat1* and *lkzf3* promoters, respectively.

Given the positive correlation between Aiolos and STAT1 expression, we performed an in silico analysis for the core IkZF factor DNA-binding motif "GGGAA" at the *Stat1* locus and found several predicted binding sites at the promoter region. Subsequent examination of publicly available Aiolos ChIP-seq data (GSM5106065) revealed a region of Aiolos enrichment at the *Stat1* promoter (Figure 8A) (23). These findings suggested that *Stat1* could be a direct target gene of

Aiolos in T_H1 cells. Since IkZF factors are known regulators of chromatin structure, we examined previously published Assay for Transposase-Accessible Chromatin (ATAC)-seq data (GSE203064) from WT and *Ikzf3^{-/-}* cells cultured under T_H1 conditions. Indeed, in the absence of Aiolos, we observed a significant decrease in chromatin accessibility at the *Stat1* promoter (Figure 8A) (28).

To test whether Aiolos could regulate Stat1 promoter activity, we created a Stat1 289 promoter-reporter construct encompassing the Aiolos-enriched region. We then overexpressed 290 with WT Aiolos or an Aiolos DNA-binding mutant (Aiolos^{DBM}) containing point mutations in the first 291 292 two N-terminal zinc finger domains, rendering this domain non-functional (Figure 8B) (18). We observed a significant increase in Stat1 promoter activity upon overexpression of WT Aiolos, and 293 this induction was lost upon overexpression of Aiolos^{DBM} (Figure 8C). These findings indicate that 294 295 Aiolos is capable of inducing Stat1 promoter activity and that the DNA-binding domain is required. 296 Lastly, to determine whether STAT1 may reciprocally regulate Aiolos expression, we examined publicly available ChIP-seq data for STAT1 (GSM994528) and previously published 297 298 ATAC-seq data (GSE203064) from WT T_H1 cells. Indeed, we identified a potential region of 299 STAT1 enrichment at the *lkzf3* promoter, which correlated with a region of accessible chromatin 300 in T_{H1} cells (Figure 8D) (28, 56). Further, ChIP analysis of IFNy-treated WT T_{H1} cells revealed that STAT1 was enriched at the *lkzf3* promoter relative to an upstream control region (Figure 8E). 301 Collectively, these data support the existence of a positive feedback loop between Aiolos and 302 303 STAT1 through which IFNy/STAT1 signaling and CXCR3 expression are regulated.

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310 Discussion

311 Aiolos has long been implicated in lymphoid cell development (16, 19-22). More recently, Aiolos has emerged as a regulator of cytokine signaling pathways and effector programs in innate and 312 adaptive lymphocytes (17, 18, 27). Here, we have identified Aiolos as a positive regulator of 313 314 IFNy/STAT1 signaling and a driver of CXCR3 expression in T_H1 cells. Our observations are 315 consistent with findings from earlier studies indicating that IFNy and STAT1 are critical for induction of CXCR3 on CD4⁺ T cells (44, 52). The work presented here expands upon those 316 findings by uncovering Aiolos as a positive transcriptional regulator of STAT1, and that 317 318 IFNy/STAT1 subsequently promotes Aiolos expression through a feed-forward amplification loop. In contrast to STAT1, we found that expression of both tyrosine-phosphorylated and bulk 319 320 STAT4 protein was elevated in Aiolos-deficient cells. Relatedly, others have reported an inverse 321 correlation between activation of STAT4 and overall levels of STAT1 in Natural Killer (NK) cells 322 (59). We also observed that Aiolos deficiency resulted in reduced expression of JAK2, a kinase of the IL-12/STAT4 signaling pathway. One explanation for this apparent contradiction may be 323 324 that alterations in expression of other cytokine signaling pathway components offsets reduced 325 JAK2 levels. For example, we have previously shown that Aiolos deficiency allows for enhanced 326 IL-2 sensitivity, and it has been established that IL-2 induces expression of *II12rb2* (28, 60). 327 Previous work has demonstrated that TYK2, another non-receptor tyrosine kinase involved in IL-12/STAT4 signaling is critical for STAT4-mediated IFNy expression (61). Future studies are 328 329 needed to determine whether TYK2 activation may contribute to sustained IL-12/STAT4 signaling 330 despite the loss of JAK2 in the absence of Aiolos.

We found that adoptively transferred Aiolos-deficient cells do not exhibit altered migration to the lungs, despite reduced CXCR3 expression. In addition to CXCR3, the chemokine receptor CCR5 is also known to enable T_H1 cell migration, and previous work has demonstrated that IL-12/STAT4 signaling selectively upregulates CCR5 (44, 62, 63). Indeed, we find elevated *Ccr5* transcript in Aiolos-deficient cells suggesting that it may compensate for reduced CXCR3

expression. Finally, given that both STAT1 and STAT4 positively impact T-bet expression, the counter-regulatory nature of Aiolos on IFN γ /STAT1 and IL-12/STAT4 signaling may help to explain the inconsistent alterations in T-bet expression that we observed in the absence of Aiolos. Our finding that CXCR3 and T-bet expression do not consistently correlate further supports the conclusion that another Aiolos-dependent factor (i.e. STAT1) is responsible for regulating CXCR3 expression. Indeed, STAT1 has been shown to control T_H1 trafficking to the lung through a T-betindependent mechanism (64).

IFNy/STAT1 signaling operates across multiple cell types suggesting that the regulatory 343 344 mechanisms established here may extend to additional Aiolos-expressing immune cells, such as Innate Lymphoid Cells (ILCs), CD8⁺ T cells, and B cells (16, 65-73). It is also possible that Aiolos 345 could impact other STAT1-driven cytokine pathways (i.e. IFN $\alpha/\beta/\lambda$, IL-27, IL-6) (74-76). For 346 347 example, a recent study indicated that Aiolos promotes the proliferation and survival of HIV-1 348 infected cells, which was associated with the upregulation of genes involved in T cell migration and type I IFN responses (77). Work by others in CD8⁺ T cells has shown that reduced levels of 349 350 STAT1, maintained by STAT4, are required for overcoming the anti-proliferative effects of type I IFN during viral infection (59). This is just one example suggesting that Aiolos-dependent 351 352 alterations in STAT1 expression could impact other signaling pathways.

Studies in humans have described IKZF3 missense mutations that compromise the DNA-353 binding domain and result in primary immunodeficiency and inborn errors of immunity (23-26). 354 Two such mutations, termed G159R and N160S, have been reported in humans and are 355 356 associated with increased susceptibility to infections and dysregulated immune responses. The IKZF3 G159R mutation has been associated with B cell deficiency, B cell malignancy, and 357 358 abnormal T cell differentiation (23). Similarly, the *IKZF3* N160S mutation has been shown to affect 359 both B and T cell populations, leading to increased susceptibility to infection. With regard to T 360 cells, the *IKZF3* N160S mutation results in decreased T_{FH} , $T_{H}1$, and memory T cell populations (25). Thus, our finding that Aiolos regulates IFNy/STAT1 signaling may provide at least some 361

mechanistic explanation for these observed disruptions in humans. Overall, these findings underscore the clinical consequences arising from genetic defects in Aiolos function, notably including compromised T_H1 responses.

While chemokine receptor expression is normally restricted to lymphocyte populations, 365 366 many cancers are known to aberrantly express these receptors and consequently acquire 367 migratory (metastatic) abilities, a phenomenon termed 'lymphocyte mimicry' (78, 79). Aiolos has been linked to metastatic lung cancer through the induction of such pathways, including 368 369 expression of CXCR4 (80-82). Of note, previous work from our lab has demonstrated that Aiolos 370 promotes CXCR5 expression in T_{FH} cells, which enables their migration into B cell follicles (18, 371 28). More recent work has demonstrated that T and B cells homozygous for an Aiolos missense 372 mutation exhibit impaired homing into lymph nodes primarily due to low CD62L expression (83). 373 These studies, alongside our observation of disrupted CXCR3 expression and altered transcript 374 levels for multiple cell adhesion molecules, integrin subunits, and chemokine receptors in Aiolos-375 deficient T_H1 cells, suggest that Aiolos drives a larger immune cell migratory program, which 376 ultimately requires further investigation.

Finally, JAK-STAT signaling pathways have long been the target of different therapeutics 377 378 due to demonstrated roles in autoimmune diseases and hematological malignancies (6-9, 84-88). 379 Similarly, Aiolos has been targeted therapeutically with lenalidomide, an immunomodulatory drug used to treat multiple myeloma and various lymphomas (89-92). Lenalidomide has also been 380 381 shown to enhance the cytotoxic activity of CAR T cells against solid tumors, which is consistent 382 with previous work showing that Aiolos suppresses T cell cytotoxic function (28, 93-95). However, 383 given the current study, it is reasonable to postulate that loss of Aiolos could also result in altered 384 lymphoid migratory patterns during immune responses to infection or cancer, which would be consistent with findings in human patients harboring Aiolos missense mutations (23-26). Hence, 385 386 therapeutics that specifically target Aiolos may have disease-specific advantages and disadvantages, presenting a potential paradox. Ultimately, future studies will be required to 387

determine the full extent of effects of Aiolos on immune cell programming, including its impact on
 STAT1-dependent signaling pathways.

390

391 Methods

392 Sex as a biological variable

393 Germline knockout influenza virus infection studies and all in vitro experiments utilized both male 394 and female mice to avoid unintentional sex bias. Similar findings are reported for both sexes. For 395 adoptive transfer studies, only male donor and recipient mice were utilized due to the Y-linked 396 nature of the OT-II transgene.

397

398 Mouse strains

399 Wild-type CD45.1 (JAX stock #002014) and CD45.2 C57BL/6J (JAX stock #000664) mice were 400 originally obtained from the Jackson Laboratory. Aiolos-deficient (*lkzf3^{-/-}*) mice were originally obtained from Riken BRC and were backcrossed onto the CD45.2 C57BL/6J Jackson background 401 402 for more than 10 generations. OT-II mice (JAX stock #004194), with the transgene located on the Y-chromosome, were originally generated by the Carbone laboratory (96) and were a generous 403 404 gift from Dr. Haitao Wen (The Ohio State University, Columbus, OH). For adoptive transfer studies, *lkzf3^{-/-}* mice were crossed to OT-II mice to generate *lkzf3^{-/-}*-OT-II mice. For all experiments 405 and replicates, individual mice were age- and sex-matched. 406

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408 CD4⁺ T cell isolation and culture

Naïve CD4⁺ T cells were isolated from the spleens and lymph nodes of 5-8-week-old mice using the BioLegend Mojo Sort naïve CD4⁺ T cell isolation kit according to the manufacturer's recommendations. For in vitro polarization of T_H1 cell populations, naïve CD4⁺ T cells were plated at a density of 300,000 cells/well in complete IMDM (IMDM [Life Technologies], 10% FBS [26140079, Life Technologies], 1% penicillin/streptomycin [Life Technologies], and 0.05% (50 μ M) 414 2-mercaptoethanol [Sigma-Aldrich]). Plates were coated with anti-CD3 (clone 145-2C11; 5 µg/mL; 415 BD Biosciences) and anti-CD28 (clone 37.51; 2 µg/mL; BD Biosciences) overnight and washed twice with PBS prior to the addition of cells in complete IMDM. Upon plating, cells were cultured 416 417 in the presence of IL-4 neutralizing antibody (clone 11B11; 5 µg/mL; BioLegend) and the T_H1-418 polarizing cytokine rmIL-12 (5 ng/mL; R&D) for 72 hours prior to analysis or expansion. For 419 experiments in which IFNy was neutralized, cells were also cultured in the presence of α -IFNy 420 antibody (clone XMG1.2; 10 µg/mL; BioLegend) for 72 hours. For expansion of cells on day 3 into 421 resting conditions, cells were plated at 500,000 cells/well in complete IMDM with the addition of 422 fresh IL-4 neutralizing antibody (clone 11B11; 5 µg/mL; BioLegend), rhIL-2 (250 U/ml; Peprotech), and either fresh rmIL-12 (5 ng/mL; R&D) or rmIFNy (50 ng/mL; Peprotech), as noted. Cells were 423 cultured for an additional 48 hours prior to harvesting for analysis on day 5. For experiments in 424 425 which cells were cultured in the presence of rmIFNy, fresh rmIFNy (50 ng/ml; Peprotech) was 426 added 1 hour prior to harvest.

427

428 **RNA isolation and qRT-PCR**

Total RNA was isolated from the cell populations described above using the Macherey-Nagel Nucleospin RNA Isolation kit according to the manufacturer's guidelines. cDNA was generated using the Superscript IV First Strand Synthesis System (Thermo Fisher Scientific). qRT-PCR reactions were performed using the SYBR Select Mastermix for CFX (Thermo Fisher Scientific) with 10 ng cDNA per reaction and primers for the appropriate genes (Supplemental Table 1). All qRT-PCR reactions were performed on the CFX Connect (BioRad). Data were normalized to *Rps18* and are presented as relative to the WT control sample.

436

437 **RNA sequencing analysis**

Published RNA-seq data (GSE203065) from WT and *lkzf3^{-/-}*T_H1 cells was analyzed as previously reported (28). Briefly, naïve CD4⁺ T cells were cultured under T_H1-polarizing conditions for 3 days.

440 Total RNA was isolated using the Macherey-Nagel Nucleospin RNA Isolation kit according to the manufacturer's guidelines. Samples were provided to Azenta Life Sciences for polyA selection, 441 library preparation, sequencing, and DESeg2 analysis (3 biological replicates per genotype from 442 443 3 independent experiments). Genes with an adjusted p < 0.05 were considered differentially 444 expressed. Heatmap generation and clustering (by Euclidean distance) were performed using 445 normalized log2 from DESeq2 analysis Morpheus software counts and the (https://software.broadinstitute.org/morpheus/). Volcano plots 446 were generated using log₁₀(adjusted p value) and log2 fold change values from DESeg2 analysis and VolcaNoseR 447 software (https://huygens.science.uva.nl/VolcaNoseR/) (97). 448

449

450 ATAC-seq analysis

451 Published ATAC-seq data (GSE203064) from WT and *lkzf3^{-/-}* cells cultured under T_H1 conditions 452 was analyzed as previously described (28, 67). Briefly, 5×10^4 cells with greater than 95% viability were processed with the Illumina Nextera DNA Library Preparation Kit according to the 453 454 manufacturer's instructions. Resultant sequences were trimmed and aligned to mm10 using 455 Bowtie2. All subsequent analyses were performed using the indicated tools in Galaxy 456 (usegalaxy.org). Samples were filtered by read quality (>30), as well as to remove duplicates and 457 mitochondrial reads. Statistically significant peaks were identified using MACS2 callpeak. DiffBind was used to identify regions of significant differential accessibility between WT and $lkzf3^{-1}$ 458 samples. Regions with adjusted p < 0.05 were considered statistically significant. CPM-459 460 normalized tracks were visualized using Integrative Genomics Viewer (IGV) versions 2.12.3 and 2.18.2. 461

462

463 **Immunoblot analysis**

Cells were harvested, counted, and lysed in 1X SDS loading dye, (50 mM Tris [pH 6.8], 100 mM
DTT, 2% SDS, 0.1% bromophenol blue, 10% glycerol) and boiled for 15 minutes. Equal protein

466 lysate amounts were loaded based on cell counts. Lysates were separated via SDS-PAGE on 467 10% Bis-Tris Bolt gels (Thermo Fisher Scientific) and then transferred onto a 0.45 µm nitrocellulose membrane. Following transfer, nitrocellulose membranes were blocked with 2% 468 469 non-fat dry milk in 1X TBST (10 mM Tris [pH 8.0], 150 mM NaCl, 0.05% Tween-20). The following 470 antibodies were used to detect proteins: α -JAK2 (1:1,000; #3230, Cell Signaling Technology), α pY-STAT4 (1:1,000; #5267, Cell Signaling Technology), α-STAT4 (1:1,000; #2653S, Cell 471 Signaling Technology), α-pY-STAT1 (1:1,000; #9167S, Cell Signaling Technology), α-STAT1 472 (1:1,000; sc-417, Santa Cruz Biotechnology), α-Aiolos (1:20,000; #39293, Active Motif), α-β-actin-473 474 HRP (1:15,000; A00730, GenScript), goat α-mouse (1:5,000; 115-035-174, Jackson Immunoresearch), and mouse α -rabbit (1:5,000-1:10,000; sc-2357, Santa Cruz Biotechnology). 475 476 Immunoblot bands were quantified using ImageJ as previously described (27). For each protein 477 row, the largest band was framed, and the mean gray value was measured using the same frame 478 across the row. Background measurements were taken with the same frame measuring the area 479 above or below bands in the image. Pixel densities were inverted, background values were 480 subtracted from sample and control bands, and a ratio of net protein bands to net loading control 481 bands was calculated for protein quantification relative to the WT sample.

482

483 Influenza virus infection and tissue preparation

Influenza A virus strain A/PR/8/34 (H1N1, "PR8") and OVA₃₂₃₋₃₃₉-expressing PR8 ("PR8-OVA") 484 485 were propagated in 10-day-old specific pathogen free embryonated chicken eggs (Charles River 486 Laboratories) and titered on MDCK cells (BEI Resources, NIAID, NIH: Kidney [Canine], Working Cell Bank, cat. # NR-2628). Mice between 8-12 weeks of age were infected intranasally with 30 487 plaque forming units (PFU) of PR8. After 8 days, mLN, spleen, and lungs were harvested as 488 489 previously described (28). For mLN and spleen, single-cell suspensions were generated in tissue 490 processing media (IMDM + 4% FBS) by passing the tissue through a 100 µm nylon mesh strainer 491 followed by erythrocyte lysis via a 3-minute incubation at room temperature in 0.84% NH₄Cl. For 492 lungs, single cell suspensions were generated by incubating whole lung tissue in HBSS (Gibco) supplemented with 1.3 mM EDTA for 30 minutes at 37°C. Following this, lungs were processed 493 in media (RPMI + 4% FBS) supplemented with Collagenase IV using a gentleMACS Dissociator 494 495 (Miltenyi Biotech) for 30 minutes according to the manufacturer's instructions. Samples were then 496 filtered through a 40 µm nylon mesh strainer and subsequently centrifuged with a Percoll density 497 gradient to isolate the mononuclear layer. Erythrocyte lysis was performed as previously described. For all tissues, cells were washed in FACS buffer (PBS + 4% FBS) prior to staining for 498 flow cytometry. For adoptive transfer studies, naïve CD45.2⁺ OT-II CD4⁺ T cells were purified from 499 WT-OT-II or *lkzf3^{-/-}*-OT-II mice using negative selection as described above. Cells were washed 500 and resuspended in sterile 1X PBS and transferred retro-orbitally (5 × 10⁵ cells/animal) into WT 501 CD45.1⁺ recipient mice that were anesthetized with isoflurane. After 24 hours, mice were infected 502 503 intranasally with 40 PFU of PR8-OVA.

504

505 Flow cytometry

For analysis of influenza nucleoprotein (NP)-specific CD4⁺ T cells in germline *lkzf3^{-/-}* animals, cells 506 were first incubated for at least 5 minutes at 4°C with TruStain FcX (anti-mouse CD16/32) Fc 507 block (clone 93; 101320; BioLegend). Samples were then stained with IA^b NP₃₁₁₋₃₂₅ MHC class II 508 509 tetramer (1:100; NIH Tetramer Core Facility) in the presence of Fc block for 1 hour at room temperature protected from light. Extracellular markers were stained in the presence of Fc block 510 511 for 30 minutes at 4°C protected from light using the following antibodies: anti-CD4 (PE/Cy7; 1:300; 512 clone GK1.5; BD Biosciences, cat. # 563933), anti-CD4 (APC; 1:300; clone GK1.5; BioLegend, cat. # 100412), anti-CXCR3 (PE; 1:300; clone CXCR3-173; BioLegend, cat. # 126505), anti-513 CXCR3 (BV421; 1:300; clone CXCR3-173; BioLegend, cat. # 126529), anti-CD44 (V450; 1:300; 514 clone IM7; BD Biosciences, cat. # 560452), anti-CD44 (FITC; 1:300; clone IM7; Thermo Fisher 515 516 Scientific, cat. # 553133), anti-CD62L (APC-efluor780; 1:300; clone MEL-14; Thermo Fisher Scientific, cat. # 47-0621-82), anti-CD45.1 (BV421; 1:300; clone A20; BioLegend, cat. # 110732) 517

518 anti-CD45.2 (APC; 1:300; clone 104; BioLegend, cat. # 109814). At the same time, cells were 519 stained with Ghost viability dye (V510; 1:400; Tonbo Biociences, cat. # 13-0870-T100). Cells were 520 then washed twice with FACS buffer prior to intracellular staining. For intracellular staining, cells were fixed and permeabilized using the eBioscience Foxp3 transcription factor staining kit 521 522 (Thermo Fisher Scientific, cat. # 00-5523-00) for 30 minutes or overnight at 4°C. After fixation, 523 samples were stained with the following antibodies in 1X eBioscience permeabilization buffer (Thermo Fisher Scientific) for 30 minutes at room temperature protected from light: anti-T-bet 524 (PerCP-Cy5.5; 1:100; clone 4B10; BioLegend, cat. # 644806) and anti-Aiolos (AF647; 1:100; 525 526 clone; S48-791; BD Biosciences, cat. # 565265). Cells were washed twice with 1X permeabilization buffer and resuspended in FACS buffer for analysis. Samples were run on a BD 527 FACS Canto II flow cytometer and analyzed using FlowJo software (version 10.8.1). 528 529 Representative gating strategies can be found in Supplemental Figures 5-7.

530

531 Promoter-reporter assay

532 A Stat1 promoter-reporter construct (pGL3-Stat1) was generated by cloning the regulatory region of Stat1 (positions – 479 to 0 bp) into the pGL3-Promoter vector (Promega) (Supplemental Table 533 534 2). Aiolos contains four N-terminal zinc finger (ZF) domains that mediate its DNA-binding capability. Of these four zinc fingers, ZF2 and ZF3 are required for DNA binding, whereas ZF1 535 and ZF4 are responsible for regulating sequence specificity (16-20). Expression vectors for WT 536 Aiolos and an Aiolos DNA-binding mutant (Aiolos^{DBM}) were constructed as previously described 537 538 (18). Briefly, two cystine residues in both ZF1 and ZF2 of Aiolos were mutated to alanine residues via site-directed mutagenesis, rendering the Aiolos DNA-binding domain non-functional. The EL4 539 murine T cell lymphoma line (TIB-39) was acquired from the American Type Culture Collection 540 541 (ATCC) and maintained in complete RPMI (RPMI-1640, 10% FBS [26140079, Life Technologies], 542 1% penicillin/streptomycin [Life Technologies]). EL4 T cell transfections were performed using the Lonza 4D nucleofection system (program CM-120, buffer SF). EL4 cells were nucleofected with 543

expression vectors for WT Aiolos, Aiolos^{DBM}, or an empty vector control in conjunction with pGL3-*Stat1* and an SV40-*Renilla* vector as a control for transfection efficiency. After 22-24 hours of
recovery, samples were harvested, and luciferase expression was analyzed using the DualLuciferase Reporter Assay System (Promega) according to the manufacturer's instructions.
Abundance of overexpressed proteins was assessed via immunoblot using an antibody against
the V5 epitope tag (Thermo Fisher Scientific, cat. # R960-25).

550

551 Chromatin Immunoprecipitation (ChIP)

552 ChIP assays were performed as described previously (98). In brief, chromatin was harvested from 553 in vitro-generated T_H 1-like cells treated with IFN γ . Chromatin was incubated with antibodies 554 against STAT1 (ThermoFisher, cat. # 10144-2-AP; 7 µg per IP) or an IgG control (Abcam, cat. # 555 ab6709; 7 µg per IP, matched to experimental antibody), and the precipitated DNA was analyzed 556 by qPCR with gene-specific primers (Supplemental Table 3). Samples were normalized to a total 557 input DNA control, and percent enrichment was divided by IgG values. The final value represents 558 the percent enrichment fold change relative to the IgG control.

559

560 Software summary

Data were collected using the following open-source or commercially available software 561 programs: BD FACSDiva (version 8.0.2), BioRad Image Lab (version 6.0.1, build 34), BioRad 562 563 CFX Manager (version 3.1). Analyses and/or manuscript preparation were conducted using BD 564 FlowJo (version 10.8.1), and open-source software, including tools available on Integrative 565 Genomics Viewer (versions 2.12.3 and 2.18.2), Morpheus VolcaNoseR 566 (https://software.broadinstitute.org/morpheus), (https://huygens.science.uva.nl/VolcaNoseR/), and (usegalaxy.org). BioRender 567 Galaxy 568 (https://biorender.com/) was used to create schematics and the graphical abstract under licenses to Leonard, M. (2024) (Supplemental Table 4). All statistical analyses were performed using 569

570 GraphPad Prism software (version 10). Data preparation for this manuscript did not require the 571 use of custom code or software.

572

573 Statistics and reproducibility

All statistical analyses were performed using GraphPad Prism software (version 10). The ROUT method (Q = 1%) was used for identifying outliers. For single comparisons, two-tailed unpaired Student's *t* test was performed. For multiple comparisons, one-way ANOVA with Tukey's multiple comparisons test was performed. Error bars indicate the standard error of the mean (SEM). The *p* values <0.05 were considered statistically significant.

579

580 Study approval

This study complies with all ethical regulations defined by the Institutional Animal Care and Use Committee (IACUC) and Institutional Biosafety Committee (IBC) of The Ohio State University in Columbus, OH (IACUC #: 2019A00000107-R1). Animals were housed in the University Laboratory Animal Resources (ULAR) Health Sciences complex at The Ohio State University in rodent barrier housing utilizing individually ventilated caging systems. All animals used in this study were humanely euthanized via CO₂ inhalation.

587

588 Data availability

Published RNA-seq (GSE203065) and ATAC-seq (GSE203064) data sets were analyzed and used in this study. The following publicly available ChIP-seq data were obtained from ChIP Atlas (https://chip-atlas.org/) for use in this study: STAT1 (GSM994528), STAT4 (GSM550303), T-bet (GSM836124), Aiolos (GSM5106065). Values for all data points in graphs are reported in the Supporting Data Values file.

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595 Author Contributions

596 M.R.L. assisted with the design of the study, performed experiments, analyzed data, and 597 wrote the manuscript. D.M.J., K.A.R., S.P., J.A.T., and R.T.W. assisted with experiments and data 598 analysis. J.S.Y. provided reagents for influenza virus infection experiments. K.J.O. supervised the 599 research, designed the study, analyzed data, and edited the manuscript.

600

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900 Figure 1. CXCR3 expression is reduced on Aiolos-deficient T_H1 cells. A) Published RNA-seq 901 data (GSE203065) from in vitro-generated WT and *lkzf3^{-/-}*T_H1 cells was assessed for differentially 902 expressed genes (DEGs). A volcano plot displays gene expression changes at day 3. Genes are color-coded: no significant changes (gray), upregulated genes with >1.5-fold change with p < 0.05903 904 (red), downregulated genes with >1.5-fold change with p < 0.05 (blue), and selected genes of interest (turquoise). **B)** Schematic of $T_{H}1$ cell culturing system. Naïve CD4⁺ T cells were stimulated 905 906 with α -CD3/CD28 under T_H1 polarizing conditions (IL-12, α -IL-4). On day 3, cells were harvested 907 or removed from stimulation and placed into fresh T_H1 conditions with IL-2 for an additional 2 days. C) At day 3, transcript analysis was performed via gRT-PCR. Transcript was normalized to 908 909 *Rps18* and presented as fold change compared to WT control (n = 10 biological replicates from 10 independent experiments, mean \pm SEM; *****p*<0.0001, two-tailed unpaired Student's *t* test). 910 911 D) Representative flow cytometric analysis for CXCR3 on day 3 T_H1 cells. Data displayed as median fluorescence intensity (MFI) fold change compared to WT controls (n = 6 biological 912 913 replicates from 6 independent experiments, mean ± SEM; ***p<0.001, two-tailed unpaired 914 Student's t test). E) At day 5, transcript analysis was performed as in 'C' (n = 9 biological replicates from 9 independent experiments, mean \pm SEM; **** p<0.0001, two-tailed unpaired Student's t 915 916 test). Note: Cxcr3 and Ikzf3 transcript data presented here are the same as in Figure 5B. F) Representative flow cytometric analysis for CXCR3 on day 5 T_H1 cells. Data displayed as MFI 917 918 fold change compared to WT controls (n = 5 biological replicates from 5 independent experiments, 919 mean ± SEM; **** p<0.0001, two-tailed unpaired Student's *t* test).

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Figure 2. CXCR3 expression is reduced on Aiolos-deficient CD4⁺ T cells responding to IAV
infection. A) Schematic of murine model of IAV infection. WT or *lkzf3^{-/-}* mice were infected
intranasally with 30 PFU of IAV (A/PR/8/34; "PR8"). After 8 days, mediastinal lymph nodes (mLN)

and lungs were harvested and stained for flow cytometric analysis. Fluorochrome-labeled MHC II tetramers were used to identify IAV nucleoprotein (NP)-specific CD4⁺ T cells. **B)** Representative flow cytometric analysis for CXCR3 expression in NP-specific CD4⁺ T cells isolated from the mLN of WT or *lkzf3^{-/-}* mice. Data are compiled from 4 independent experiments and displayed as percent positive for CXCR3. **C)** Representative histogram overlay for CXCR3. Data are displayed as MFI fold change compared to WT controls (*n* = 16 for WT and *n* = 15 for *lkzf3^{-/-}*, mean ± SEM; ******p*<0.0001, two-tailed unpaired Student's *t* test).

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Figure 3. CXCR3 expression is reduced on Aiolos-deficient CD4⁺ T cells in a cell-intrinsic 932 manner. A) Schematic of adoptive transfer system. Naïve CD4⁺T cells were harvested from the 933 mLN of WT-OT-II or *lkzf3^{-/-}*-OT-II mice. 500,000 cells/animal were adoptively transferred into 934 935 CD45.1⁺ recipients. Recipient mice were then infected with 40 PFU of OVA₃₂₃₋₃₃₉-expressing 936 A/PR/8/34 ("PR8-OVA") influenza virus 24 hours post-transfer. 8 days post-infection, mLN was harvested and viable CD45.2+CD4+CD62L-CD44+ (antigen-specific, donor effector) cells were 937 938 analyzed via flow cytometry. B) Representative flow cytometric analysis for CXCR3 expression in CD45.2⁺CD4⁺CD62L⁻CD44⁺ cells in the mLN. Data are compiled from 3 independent experiments 939 940 and displayed as percent positive for CXCR3. C) Representative histogram overlay for CXCR3. Data are displayed as MFI fold change compared to WT-OT-II control cells (n = 13, mean \pm SEM; 941 ****p*<0.001, *****p*<0.0001, two-tailed unpaired Student's *t* test). 942

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Figure 4. Aiolos-deficient $T_{H}1$ cells exhibit altered expression of components of IFNY/STAT1 and IL-12/STAT4 signaling pathways. A) Publicly available Chromatin Immunoprecipitation sequencing (ChIP-seq) data for STAT1 (GSM994528), STAT4 (GSM550303), and T-bet (GSM836124) was examined at *Cxcr3*. Sequencing tracks were viewed using the Integrative Genomics Viewer (IGV). Regulatory regions of interest with transcription factor enrichment are indicated by the blue boxes. **B)** Published RNA-seq data (GSE203065)

from in vitro-generated WT and $lkzf3^{-1}T_H1$ cells was analyzed for differentially expressed genes 950 951 (DEGs). A heatmap of DEGs associated with IFNy/STAT1 and IL-12/STAT4 signaling in T_{H1} cells 952 is shown, as well as additional genes involved in both pathways and T helper cell differentiation. Gene names color-coded in blue are downregulated in *lkzf3^{-/-}* T_H1 cells. Note: *Cxcr3* transcript 953 954 data presented here are the same as in Supplemental Figure 3D. C) Schematic of proposed 955 model in which Aiolos may regulate CXCR3 via impacts on components of the IFNy/STAT1 and IL-12/STAT4 cytokine signaling pathways. The downward arrows in blue indicate genes that are 956 957 downregulated in the absence of Aiolos.

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Figure 5. IFNy/STAT1 signaling, but not IL-12/STAT4, is diminished in IL-12-treated Aiolos-959 deficient T_H1 cells. A) Schematic of culturing system. Naïve CD4⁺ T cells were stimulated with 960 961 α -CD3/CD28 under T_H1 polarizing conditions (IL-12, α -IL-4). On day 3, cells were removed from 962 stimulation and placed back into fresh T_H1 polarizing conditions (IL-12, α -IL-4) with IL-2 for an additional 2 days. B) At day 5, transcript analysis was performed via qRT-PCR. Transcript was 963 964 normalized to Rps18 and presented as fold change compared to WT control (n = 8-9 biological replicates from 8-9 independent experiments. Data are presented as mean \pm SEM; ***p<0.001, 965 966 ****p<0.0001, two-tailed unpaired Student's t test). Note: Cxcr3 and lkzf3 transcript data 967 presented here are the same as in Figure 1E. C-D) An immunoblot was performed to assess the relative abundance of the indicated proteins. β -actin serves as a loading control (n = 5-7968 independent experiments, mean ± SEM; *p<0.05, ***p<0.001, ****p<0.0001, two-tailed unpaired 969 970 Student's *t* test).

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Figure 6. IFNγ/STAT1 signaling is compromised in IFNγ-treated Aiolos-deficient T_H1 cells. A) Schematic of culturing system. Naïve CD4⁺ T cells were stimulated with α-CD3/CD28 and cultured under T_H1 polarizing conditions (IL-12, α-IL-4). On day 3, cells were removed from stimulation and given IFNγ, α-IL-4, and IL-2 for an additional 2 days. **B)** At day 5, transcript

976 analysis was performed via gRT-PCR. Transcript was normalized to Rps18 and presented as fold 977 change compared to WT control (n = 4 biological replicates from 4 independent experiments, mean \pm SEM; ***p*<0.01, ****p*<0.001, *****p*<0.0001, two-tailed unpaired Student's *t* test). **C)** 978 979 Representative flow cytometric analysis for CXCR3 on IFNy-treated $T_{H}1$ cells at day 5. Data are 980 displayed as MFI fold change compared to WT controls (n = 3 biological replicates from 3 independent experiments, mean \pm SEM; **p<0.01, two-tailed unpaired Student's t test). **D)** An 981 immunoblot was performed to assess the relative abundance of the indicated proteins. β-actin 982 serves as a loading control (n = 4 independent experiments, mean \pm SEM; *p < 0.05, ***p < 0.001, 983 two-tailed unpaired Student's t test). E) ChIP assays were performed to assess STAT1 984 association with Cxcr3 in WT and Ikzf3^{-/-}T_H1 cells. Publicly available ChIP-seq data for STAT1 985 (GSM994528) was examined to identify potential regions of STAT1 enrichment. Sequencing 986 987 tracks were viewed using IGV and regulatory regions of interest are indicated by blue boxes. 988 Approximate ChIP primer locations at the Cxcr3 promoter ("prom.") and 3' enhancer ("enhc.") are indicated with gray arrows. F) The indicated regions were analyzed for STAT1 enrichment. Data 989 990 were normalized to total input. Percent enrichment relative to input was divided by IgG, and data are presented as fold change relative to IgG. (n = 4 biological replicates from 4 independent 991 992 experiments, mean ± SEM; **p<0.01, ***p<0.001, one-way ANOVA with Tukey's multiple 993 comparisons test).

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Figure 7. IFNγ/STAT1 signaling induces Aiolos expression. A) Schematic of culturing system. WT naïve CD4⁺ T cells were stimulated with α-CD3/CD28 under T_H1 polarizing conditions (IL-12, α-IL-4). Some cells were also treated with α-IFNγ to inhibit IFNγ/STAT1 signaling. **B)** At day 3, transcript analysis was performed via qRT-PCR. Transcript was normalized to *Rps18* and presented as fold change compared to WT control (n = 4 biological replicates from 4 independent experiments, mean ± SEM; **p<0.01, ****p<0.0001, two-tailed unpaired Student's *t* test). **C)** Representative flow cytometric analysis at day 3 for CXCR3 expression on WT T_H1 cells treated 1002 with or without α -IFNy. Data are displayed as percent positive for CXCR3 (n = 3 biological 1003 replicates from 3 independent experiments, mean ± SEM; *p<0.05, two-tailed unpaired Student's 1004 t test). D) An immunoblot was performed to assess the relative abundance of the indicated 1005 proteins. β -actin serves as a loading control (n = 4 independent experiments, mean \pm SEM; 1006 *p < 0.05, **p < 0.01, ****p < 0.0001, two-tailed unpaired Student's t test). E) At day 3, transcript and 1007 flow cytometric analyses were performed for *lkzf3* and Aiolos protein expression, respectively. 1008 Flow cytometric data are displayed as MFI fold change compared to WT controls (n = 3 biological replicates from 3 independent experiments, mean ± SEM; **p<0.01, two-tailed unpaired Student's 1009 1010 t test).

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1012 Figure 8. Aiolos and STAT1 engage in reciprocal regulation. A) Publicly available ATAC-seq 1013 data (GSE203064) from WT and *lkzf3^{-/-}* T_H1 cells was assessed for alterations in chromatin 1014 accessibility at the Stat1 promoter. Publicly available ChIP-seq data for Aiolos (GSM5106065) 1015 was examined at Stat1. Sequencing tracks were viewed using IGV. The Stat1 promoter region of significant differential accessibility is indicated by a blue box ($p_{adj} = 0.0302$). A ~500 bp region 1016 1017 encompassing the indicated Aiolos DNA binding motifs within the Stat1 promoter was cloned into 1018 a reporter plasmid. B) Schematic depicting the zinc finger (ZF) domains of WT Aiolos and a DNAbinding mutant (Aiolos^{DBM}). C) EL4 T cells were transfected with a Stat1 promoter-reporter and 1019 either WT Aiolos, Aiolos^{DBM}, or empty vector control. Cells were also transfected with SV40-1020 1021 Renilla as a control for transduction efficiency. Luciferase promoter-reporter values were 1022 normalized to Renilla control and presented relative to the empty vector control. Aiolos was 1023 assessed via immunoblot with an antibody for the V5 epitope tag. β -actin serves as a loading 1024 control. Data are representative of 3 independent experiments (n = 3, mean \pm SEM; *p < 0.05, one-1025 way ANOVA with Tukey's multiple comparisons test). D) Publicly available ATAC-seq data 1026 (GSE203064) from T_H1 cells and ChIP-seq data for STAT1 (GSM994528) were viewed using IGV to identify regions of STAT1 enrichment (blue box) at *Ikzf3*. Approximate ChIP primer locations 1027

are indicated with a gray arrow. **E)** The *lkzf3* promoter ("prom.") and a negative control region ("neg. ctrl.") were analyzed for STAT1 enrichment via ChIP. Data were normalized to total input. Percent enrichment relative to input was divided by IgG, and data are presented as fold change relative to IgG. (n = 4 biological replicates from 4 independent experiments, mean \pm SEM; *p<0.05, **p<0.01, one-way ANOVA with Tukey's multiple comparisons test).

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1034 Supplemental Figure 1. Aiolos-deficient mice have reduced numbers of CD4⁺ T cells in the lungs during IAV infection. WT or *lkzf3^{-/-}* mice were infected intranasally with 30 PFU of IAV 1035 1036 (A/PR/8/34; "PR8"). After 8 days, mLN and lungs were harvested and stained for flow cytometric 1037 analysis. Fluorochrome-labeled MHC II tetramers were used to identify IAV nucleoprotein (NP)specific CD4⁺ T cells. A-B) NP-specific CD4⁺ T cells were enumerated. Cell numbers were 1038 1039 normalized to 1 x 10⁶ total events. Numbers and percentages of NP-specific CD4⁺ T cells in the 1040 mLN and lungs are displayed. Data from 4 independent experiments is shown (For cell numbers, n = 17 for WT and n = 15 for *lkzf3^{-/-}*. For percentages, n = 14 for mLN and n = 15 for lungs. Data 1041 are presented as mean \pm SEM; *****p*<0.0001, two-tailed unpaired Student's *t* test). **C-D)** Bulk 1042 1043 CD4⁺ T cells were enumerated. Cell numbers were normalized to 1 x 10⁶ total events. Numbers 1044 and percentages of bulk CD4⁺ T cells in the mLN and lungs are displayed. Representative data from 4 independent experiments shown (For cell numbers, n = 17 for WT and n = 15 for $lkzf3^{-/-}$. 1045 For percentages, n = 15 for mLN and n = 14 for lungs. Data are presented as mean \pm SEM; 1046 **** p<0.0001, two-tailed unpaired Student's t test). 1047

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1049 Supplemental Figure 2. Aiolos-deficient mice exhibit no change in T-bet expression but 1050 have decreased CXCR3 expression during IAV infection. WT or *lkzf3^{-/-}* mice were infected 1051 intranasally with 30 PFU of IAV (A/PR/8/34; "PR8"). After 8 days, mLN and spleen were harvested 1052 and stained for flow cytometric analysis. Fluorochrome-labeled MHC II tetramers were used to 1053 identify IAV nucleoprotein (NP)-specific CD4⁺ T cells in the mLN. **A)** Representative flow

1054 cytometric analysis for T-bet expression in NP-specific cells isolated from the mLN of WT or *lkzf3*⁻ 1055 ⁷ mice. Data are compiled from 4 independent experiments and displayed as MFI fold change compared to WT controls (n = 16 for WT and n = 15 for $lkzf3^{-/-}$, mean \pm SEM; two-tailed unpaired 1056 1057 Student's *t* test). **B-D)** Representative flow cytometric analyses for CXCR3 expression in bulk 1058 CD4⁺ naïve (CD62L⁺CD44⁻), central memory (CD62L⁺CD44⁺), and effector (CD62L⁻CD44⁺) T cell populations isolated from the spleens of WT or $lkzf3^{-/-}$ mice. Data are compiled from 4 independent 1059 1060 experiments and displayed as MFI fold change compared to WT controls (n = 17 for WT and n =15 for $lkzf3^{-/}$, mean ± SEM; *p<0.05, ****p<0.0001, two-tailed unpaired Student's t test). 1061

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1063 Supplemental Figure 3. CD4⁺ T cells display altered migratory programs in the absence of Aiolos. Naïve CD4⁺ T cells were harvested from the mLN and lungs of WT-OT-II or *lkzf3^{/-}*-OT-II 1064 1065 mice. 500,000 cells/animal were adoptively transferred into CD45.1⁺ recipients. Recipient mice 1066 were then infected with 40 PFU of OVA₃₂₃₋₃₃₉-expressing A/PR/8/34 ("PR8-OVA") influenza virus 24 hours post-transfer. 8 days post-infection, mLN and lungs were harvested and viable 1067 CD45.2⁺CD4⁺CD62L⁻CD44⁺ (antigen-specific, donor effector) cells were analyzed via flow 1068 1069 cytometry. A) Representative flow cytometric analysis for T-bet expression in 1070 CD45.2⁺CD4⁺CD62L⁻CD44⁺ cells in the mLN. Data are compiled from 3 independent experiments and displayed as MFI fold change compared to WT-OT-II control cells (n = 13, mean \pm SEM; 1071 ***p<0.001, two-tailed unpaired Student's t test). **B-C)** Total CD45.2⁺ cell numbers were 1072 1073 enumerated. Cell numbers were normalized to 500,000 total events. Numbers and percentages 1074 of CD45.2⁺ cells in the mLN and lungs are displayed. Data from 3 independent experiments is shown (For cell numbers, n = 14-15. For percentages, n = 13-15. Data are presented as mean ± 1075 SEM; **p<0.01, two-tailed unpaired Student's t test). **D)** Published RNA-seq data (GSE203065) 1076 from in vitro-generated WT and $lkzf3^{-7}$ T_H1 cells was analyzed for differentially expressed genes 1077 1078 (DEGs). A heatmap of DEGs associated with cell migration in $T_H 1$ cells is shown. Gene names color-coded in blue are downregulated in *lkzf3^{-/-}* T_H1 cells. Gene names color-coded in red are 1079

1080 upregulated in $lkzf3^{-}$ T_H1 cells. Note: *Cxcr*3 transcript data presented here is the same as in 1081 Figure 4B. RTK; receptor tyrosine kinase. GEF; guanine nucleotide exchange factor.

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1083 Supplemental Figure 4. IFNy/STAT1 and IL-12/STAT4 pathways are altered in Aiolos-1084 deficient T_H1 cells. Naïve CD4⁺ T cells were harvested from WT and *lkzf3^{-/-}* mice and stimulated with α -CD3/CD28 under T_H1 polarizing conditions (IL-12, α -IL-4). On day 3, cells were removed 1085 1086 from stimulation and given either 1.) IL-12, α-IL-4, and IL-2 or 2.) IFNy, α-IL-4, and IL-2 for an additional 2 days prior to harvest. A) At day 5, transcript analysis was performed on IL-12-treated 1087 1088 T_H1 cells via gRT-PCR. Transcript was normalized to *Rps18* and presented as fold change 1089 compared to WT control (n = 4-8 biological replicates from 4-8 independent experiments. Data are presented as mean ± SEM; *** p<0.001, **** p<0.0001, two-tailed unpaired Student's t test). 1090 1091 **B)** At day 5, RNA was isolated from IFNγ-treated T_H1 cells, and transcript analysis was performed 1092 as in 'A' (n = 4 biological replicates from 4 independent experiments, mean ± SEM; *p < 0.05, ***p < 0.001, two-tailed unpaired Student's t test). C) An immunoblot of IFNv-treated T_H1 cells was 1093 1094 performed to assess the relative abundance of the indicated proteins. β-actin serves as a loading 1095 control (n = 4 independent experiments, mean \pm SEM; **p<0.01, two-tailed unpaired Student's t 1096 test).

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Supplemental Figure 5. Representative flow cytometry gating strategy for mediastinal lymph node (mLN) and lungs in germline knockout IAV infection experiments. WT or *lkzf3*^{-/-} mice were infected intranasally with 30 PFU of IAV (A/PR/8/34; "PR8"). After 8 days, mLN and lungs were harvested and stained for flow cytometric analysis of CXCR3 and T-bet expression in IAV nucleoprotein (NP)-specific CD4⁺ T cells. Fluorochrome-labeled MHC II tetramers were used to identify NP-specific cells in the mLN and lungs.

Supplemental Figure 6. Representative flow cytometry gating strategy for spleen in germline knockout IAV infection experiments. WT or *lkzf3^{-/-}* mice were infected intranasally with 30 PFU of IAV (A/PR/8/34; "PR8"). After 8 days, spleen was harvested and stained for flow cytometric analysis of CXCR3 expression in bulk CD4⁺ naïve (CD62L⁺CD44⁻), central memory (CD62L⁺CD44⁺), and effector (CD62L⁻CD44⁺) T cell populations.

1111	Supplemental Figure 7. Representative flow cytometry gating strategy for adoptive transfer
1112	experiments. Naïve CD4 ⁺ T cells were harvested from the mLN of WT-OT-II or <i>lkzf3^{-/-}</i> -OT-II mice.
1113	500,000 cells/animal were adoptively transferred into CD45.1+ recipients. Recipient mice were
1114	then infected with 40 PFU of OVA ₃₂₃₋₃₃₉ -expressing A/PR/8/34 ("PR8-OVA") influenza virus 24
1115	hours post transfer. 8 days post-infection, mLN was harvested and viable CD45.2+CD4+ CD62L-
1116	CD44 ⁺ (antigen-specific, donor effector) cells were analyzed via flow cytometry for CXCR3 and
1117	T-bet expression.
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Main Figures











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1207 Supplemental Materials

1208 Supplemental Figure 1



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Supplemental Table 1. qRT-PCR primers

Gene (murine)	Forward (5'-3')	Reverse (5'-3')
Rps18	GGAGAACTCACGGAGGATGAG	CGCAGCTTGTTGTCTAGACCG
lkzf3	CCGACTGTGGAGCTGAAAAGC	CCTGCATCTTCGTCTTCATTGG
Cxcr3	CCTTGAGGTTAGTGAACGTC	GCTGGCAGGAAGGTTCTGTC
Tbx21	GTGACTGCCTACCAGAACGC	AGGGGACACTCGTATCAACAG
Stat1	GGTACAACATGCTGGTGACAGAG	CTCCCAGCATGCTCAGCTGGTC
Stat4	CCAATGGGAGCCTCTCAGTGGAG	GCAACTCCTCTGTCACCATGTG
Jak1	GCTGAGGTGGAGCTGCACCGAC	GTCCATAGAGCCATGCAGGCTG
Jak2	GGAAACTTGGAGTGGCTAAGCAG	GTGGGTTCCCCGTTCTCCTGTC
Jak3	CCTGATCTGCGACTCCAGGC	GAGAATGTAGGTGCCTGGGAG
Tyk2	GGAGCGTCGCGTGCACATCCAC	GTGGCTGGAGTCAGCAGTCAAGC
lfngr1	GTGTATGTGGAGCATAACCGGAG	CTGGAATCCAGTGTGGATACTGAG
lfngr2	GAGCAATGTATCCTGTCACG	GTCAGGCCGAGCAGCAATGCG
ll12rb1	CACGACTCGGCTCCTCATGGAC	TCTCAACGCAGCCATCACC
ll12rb2	CTTGGACGGCATCAGTGTCTGC	GACCTGGTGAGGAGCCAGCAAC

Supplemental Table 2. Promoter-reporter primers

Gene (murine)	Forward (5'-3')	Reverse (5'-3')
Stat1 prom.	GATCGGTACCGCAGGCTTGGTTGACGTCAGTG	GATCGAGCTCAGGGCGTCCCGCCTCCTTCCGCCTC
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1292 Supplemental Table 3. ChIP qPCR primers

Gene (murine)	Forward (5'-3')	Reverse (5'-3')
Cxcr3 prom.	CAGGTCTCGTGCTGCCTGCTTCTC	CTGCGGAGGGCTGGTATAGATTACC
Cxcr3 enhc.	GGGAGAAAGTGACAGTGCAG	CAGACATTAGCATGAAGCCACC
Cxcr3 ctrl.	GCCTAGGGAAGATAGTTCCTC	GGTTGAAGCAGGGAGTGGTGG
<i>lkzf3</i> prom.	GACGTCTACTTGAGAAACACCGG	CACTGACAGTTCTCAAGACCGTC
<i>lkzf3</i> ctrl.	GTGCAGCTTCCCAATAAACCTGCC	GGAACTCACCATGTAGACCAGGCTG

Figure	Citation
Graphical abstract	Leonard, M. (2024) https://BioRender.com/i42i899
Figure 1B	Leonard, M. (2024) https://BioRender.com/v70c175
Figure 2A	Leonard, M. (2024) https://BioRender.com/p22p915
Figure 3A	Leonard, M. (2024) https://BioRender.com/h43w136
Figure 4C	Leonard, M. (2024) https://BioRender.com/b87z682
Figure 5A	Leonard, M. (2024) https://BioRender.com/i17z581
Figure 6A	Leonard, M. (2024) https://BioRender.com/u39m539
Figure 7A	Leonard, M. (2024) https://BioRender.com/r22y787

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