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Cellular Importin-α3 Expression Dynamics in the Lung Regulate Antiviral Response Pathways against Influenza A Virus Infection

Graphical Abstract

Highlights

- Importin-α3 is the most abundantly expressed isoform in the mammalian lung
- Importin-α3 is highly conserved across species
- Importin-α3 is one of the major nuclear transporters of NF-κB
- Importin-α3 acts as an immune sensor of influenza A virus infections

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In Brief

Thiele et al. show that importin-α3 is one of the major nuclear transporters of NF-κB in the mammalian lung. High-level TNF-α-inducing HPAIVs inhibit importin-α3 mRNA transcription by interfering with its promoter activity. Thus, HPAIVs may evade antiviral immunity in the respiratory tract by generating a bottleneck in importin-α3 availability.
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Cellular Importin-α3 Expression Dynamics in the Lung Regulate Antiviral Response Pathways against Influenza A Virus Infection

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SUMMARY

Importin-α adaptor proteins orchestrate dynamic nuclear transport processes involved in cellular homeostasis. Here, we show that importin-α3, one of the main NF-κB transporters, is the most abundantly expressed classical nuclear transport factor in the mammalian respiratory tract. Importin-α3 promoter activity is regulated by TNF-α-induced NF-κB in a concentration-dependent manner. High-level TNF-α-inducing highly pathogenic avian influenza A viruses (HPAIVs) isolated from fatal human cases harboring human-type polymerase signatures (PB2 627K, 701N) significantly downregulate importin-α3 mRNA expression in primary lung cells. Importin-α3 depletion is restored upon back-mutating the HPAIV polymerase into an avian-type signature (PB2 627E, 701D) that can no longer induce high TNF-α levels. Importin-α3-deficient mice show reduced NF-κB-activated antiviral gene expression and increased influenza lethality. Thus, importin-α3 plays a key role in antiviral immunity against influenza. Lifting the bottleneck in importin-α3 availability in the lung might provide a new strategy to combat respiratory virus infections.

INTRODUCTION

Molecular trafficking of proteins between the cytoplasm and the nucleus is a fundamental process that is essential to maintain cellular homeostasis ( Görlich and Mattaj, 1996; Imamoto et al., 1995). Disturbances in these highly sensitive regulatory processes may result in the disequilibrium of cellular and nuclear proteins and eventually cause diseases. Importin-α proteins belong to the major nuclear transport factors in the cell shuttling various cargo proteins containing a nuclear localization signal (NLS) from the cytoplasm to the nucleus (Goldfarb et al., 2004). Importin-α proteins are acidicophilic proteins that bind as adaptors to cargo proteins containing a basic stretch of mono- or bipartite NLS motifs. This importin-α/cargo complex forms a trimeric complex with the receptor protein importin-β1, which finally translocates through the nuclear pore by the interaction of importin-β1 with nucleoporins (Hu et al., 1996; Radu et al., 1995; Rexach and Blobel, 1995). In the nucleus, the guanosine triphosphate (GTP)-bound form of Ran binds to importin-β1 and mediates the dissociation and release of the cargo protein into the nucleus (Lee et al., 2005; Matsuura and Stewart, 2005). After successful nuclear import, importin-α binds to the recycling factor cellular apoptosis susceptibility protein (CAS) that, in conjunction with RanGTP, transports the complex back into the cytoplasm where importin-α is again available for the nuclear import of other cargo proteins (Kutay et al., 1997). Importin-α proteins are highly conserved throughout evolution (Goldfarb et al., 2004). In eukaryotes, seven importin-α isoforms were described, whereas the budding yeast encodes one importin-α isoform, and flies and nematodes encode for three importin-α isoforms. However, all of these organisms encode for only one importin-β1 receptor protein. The specificity of importin-α isoforms to cargo proteins is determined by the complexity and the three-dimensional context (Friedrich et al., 2006; Goldfarb et al., 2004; Sankala et al., 2017). Moreover, importin-α proteins play a key role in cellular differentiation by regulating the transport of transcription factors.
factors through lineage-specific expression profiles in cells (Yasu­hara et al., 2007). However, little is known about the expression profiles of importin-α isoforms in organs that could give important insights into molecular mechanisms in health and disease. There is accumulating evidence that under cellular stress, such as oxidative stress, heat shock, or ultraviolet irradiation, importin-α proteins may accumulate in the nucleus and disrupt cellular homeostasis (Furuta et al., 2004; Kodina et al., 2004; Yasuda et al., 2012). In the last decade, a growing amount of evidence suggests that importin-α proteins are utilized by various viruses (e.g., influenza, Ebola, dengue, hepatitis C, and corona­viruses) to circumvent host defense mechanisms (Basler and Amarasinge, 2009; Canton et al., 2018; Fraser et al., 2014; Fries­man et al., 2007; Gabriel et al., 2011; Hudjetz and Gabri­el, 2012; Kopecky-Bromberg et al., 2007; Neufeldt et al., 2013; Pryor et al., 2007). Little evidence is available on their mode of action in specific organs under inflammatory conditions such as virus infections. In this study, we identified the anatomic expression profile of importin-α isoforms in the mammalian lung and studied their transcriptional regulation and impact on influenza-virus­induced pneumonia using cell culture models, genome-wide transcription analyses, and transgenic mouse models.

RESULTS

Importin-α3 Is the Most Abundantly Expressed Importin-α Isoform in the Mammalian Respiratory Tract

In order to analyze the expression pattern of major importin-α isoforms throughout the mammalian respiratory tract (RT), we anatomic­ally dissected the human and murine RT into an upper part defined by respiratory cells of the nasal concha (URT) and a lower part defined by bronchi and alveoli (LRT). Importin-α1, -α3, and -α5/7 isoforms (Kpna2, Kpna4, and Kpna1/6, respectively) were abundantly expressed in epithelial cells, glands, and macrophages of the human (Figure 1A) as well as the murine URT and LRT (Figure 1B). We then measured the relative mRNA expression levels of importin-α isoforms along the mammalian RT. In the human URT and LRT, importin-α3 was by far the most abundantly expressed isoform, with 4–5 times increased mRNA levels compared to importin-α1. Average expression levels of other isoforms were similar to those of importin-α1, except for importin-α7, which was the second-most abundant isoform (Figure 1C). Similarly, importin-α3 was also the highest-expressed isoform, followed by importin-α7 in the murine URT and LRT. Laser-based microdissection of epithelial cells was used to ensure homogeneous cell populations (Figure 1D). We further confirmed the exclusive abundance of importin-α3 in the murine lung, followed by importin-α7 as the second-most abundant isoform also on the protein level, using whole-organ homogenates (Figures 1E and 1F). Notably, in immortalized cells (Figures S1A–S1D) and unlike in primary cells, importin-α1 is the most abundant isoform and is therefore used as a prognostic cancer marker (Christiansen and Dyskjøt, 2013; Dahl et al., 2006; Kau et al., 2004). Thus, it is particularly important to investigate the role of importin-α isoforms either in primary cells or in an in vivo animal model. We then determined the expression of importin-α isoforms in various murine organs on the mRNA and protein levels and confirmed that the lung (Figure S1E) is unique in exhibiting exclusively high importin-α3 levels, compared to other solid organs (Figures S1F–S1K). Moreover, comparable distribution and expression levels of importin-α isoforms within the human and murine RT suggest that the mouse model may be considered adequate to study the impact and mode-of-action of these cellular factors in health and disease. Next, we directly compared relative importin-α amounts in laser-microdissected epithelial samples of the URT versus the LRT (Figures 1G–1K). For all importin-α isoforms, mRNA levels were highest in the alveolar tissue (LRT (A), up to 6 times) followed by epithelial cells of the bronchi and bronchioles (LRT [bronchial epithelium, BE]). The lowest mRNA levels were detected in epithelial cells of the nasal concha (URT). Thus, importin-α isoforms are expressed as a gradient in the RT with gradually increasing mRNA levels from the URT toward the LRT (Figure 1L). However, importin-α3 is the most abundantly expressed importin-α isoform in the mammalian RT, unlike in other solid organs.

Importin-α3 Mediates Nuclear Translocation of TNF-α-Activated NF-κB

NF-κB subunits (p50, p65) are translocated into the nucleus in an importin-α-mediated manner. Herein, particularly importin-α3 and its close relative, importin-α4, were shown to act as one of the main isoforms that imports tumor necrosis factor alpha (TNF-α)-activated p50/p65 heterodimers into the nucleus via direct interaction (Fagerlund et al., 2005). Since the mammalian lung most abundantly expresses the importin-α3 isoform, we continued to dissect the physical and functional interaction between importin-α3 and NF-κB. We could confirm that purified human importin-α3 protein directly interacts with NF-κB p50 and p65 subunits as well as its precursor p105 protein upon activation with TNF-α unlike importin-α1 that was used as a negative control (Figures 2A and 2B). Consistently, treatment of murine fibroblasts with TNF-α most significantly increased NF-κB levels in nuclear cell fractions along with importin-α3 (Figure 2C). In order to confirm that importin-α3 is required for the nuclear import of NF-κB, we generated murine embryo fibroblasts (MEFs) with a deleted importin-α3 gene (α3−/−). Nuclear fractionation analysis in TNF-α-treated α3+/− MEFs showed significantly reduced nuclear translocation of the NF-κB subunit p65 compared to wild-type (WT) control (α3+/+) MEFs (Figure 2D). Immunofluorescence analysis further confirmed that nuclear localization of NF-κB p65 was also reduced in α3−/− MEFs compared to α3+/− MEFs (Figure 2E). Conversely, NF-κB p65 accumulated in the cytoplasm of α3−/− MEFs, albeit slightly, likely due to the degradation of inefficiently transported cargo proteins (Figure 2E). The extent of reduced nuclear localization of importin-α3 was dependent on the method used, albeit the absence of importin-α3 consistently impaired efficient nuclear localization of TNF-α-induced NF-κB subunit p65. These data clearly show that importin-α3 acts as a nuclear translocator of TNF-α-activated NF-κB.

Importin-α3 Promoter Activity Is Regulated by TNF-α-Activated NF-κB

We further observed an increase in general importin-α3 protein levels upon TNF-α treatment in total cell lysates, including nuclear and cytoplasmic cell fractions of murine as well as primary
Figure 1. Importin-α Expression in the Mammalian Respiratory Tract

(A and B) Importin-α1, -α3, or -α5/7 proteins were stained (red) in human (A) or murine (B) upper (URT, nasal epithelial cells) and lower respiratory tract (LRT, bronchi and alveoli) sections by IHC-P (immunohistochemistry-paraffin protocol) and counterstained with hematoxylin. The importin-α7 antibody cross-reacts with importin-α5. 400× (URT and bronchi in A) or 1000× (alveoli in A and B) original magnification. Scale bar, 10 μm.

(C and D) Importin-α mRNA levels in the human (C) and murine (D) URT and LRT determined by qRT-PCR. Relative importin-α1 expression values were set to 1 after normalization against GAPDH (Glycerinaldehyde-3-phosphat-dehydrogenase). Each data point represents an individual sample (n = 6–12).

(E and F) Importin-α protein amounts in murine LRT. Western blot analyses using importin-α isoform-specific antibodies and GAPDH adjustment were performed to determine endogenous importin-α protein amounts (α1, α3, α5, α7) in murine organs. For relative quantification (E), standard curves of affinity-purified, FLAG-tagged importin-α proteins were used. Relative importin-α1 protein amount in murine LRT was set to 1. Data shown represent means ± SD (n = 3 biological replicates; technical replicates: n = 1–2 per organ, n = 1–2 western blot analyses). Representative western blots for the endogenous importin-α isoforms and the standard curves are shown in (F). The gaps in (F) depict cropping of the relevant bands run on the same gel.

(G–K) Comparison of importin-α mRNA expression levels between murine URT and LRT: α1 (G), α3 (H), α4 (I), α5 (J), or α7 (K). A, alveoli; BE, bronchiolar epithelium; n = 8–12.

(L) Schematic partition of the RT with increasing importin-α mRNA levels from URT to LRT.

*p < 0.05; **p < 0.01; ***p < 0.001.
airway epithelial cells (Figures 2F and 2G). These findings might suggest that importin-α3 gene expression itself could also be affected by TNF-α treatment. In order to understand how importin-α3 gene transcription is regulated, we screened the promoter region of the importin-α3 gene for the presence of potential transcription factor binding sites. We found that the promoter region of the importin-α3 gene contains a binding site for NF-κB, as shown by earlier chromatin immunoprecipitation studies (Raskato et al., 2012) (Figure 2H). Consistently, using a reporter construct encoding for renilla luciferase under the control of the importin-α3 promoter, we could show that importin-α3 gene expression is differentially affected upon TNF-α treatment in a cell-type- and dose-dependent manner (Figures 2I–2K). In clinically relevant primary human airway epithelial cells as well as in generated MEFs herein, importin-α3 promoter activity was highest upon low-dose TNF-α treatment, while high-dose TNF-α treatment resulted in a reduction of importin-α3 gene expression (Figures 2J and 2K). However, multi-passage cancerous A549 cells presented high resistance to TNF-α treatment (Figure 2L), suggesting a delayed dose-response curve. Treating MEFs with other cytokines, such as interleukin (IL)-10 or transforming growth factor β (TGF-β) did not significantly alter importin-α3 promoter activity, unlike TNF-α treatment (Figures S2A and S2B), albeit IL-10 presented a slight induction (Figure S2A). Next, we wanted to assess whether TNF-α-activated NF-κB is directly involved in regulating importin-α3 transcription. For this, we used cells deficient in the NF-κB gene essential modulator (NEMO), the regulatory subunit of the IKK (IκB kinase), which is required for NF-κB phosphorylation and thus its activation. Importin-α3 mRNA levels were reduced in TNF-α-treated NEMO+/− cells compared to untreated controls (Figure 2L). As expected, reduced importin-α3 mRNA levels also resulted in reduced importin-α3 protein expression in NEMO+/−cells treated with TNF-α as compared to untreated cells (Figure 2M). Reduced importin-α3 expression in NEMO+/−cells could be restored upon transfection of a NEMO-expressing plasmid back into NEMO−/−cells (Figure 2M). Control treatment of NEMO−/−cells with IL-6, IL-10, interferon (IFN)-γ or TGF-β showed similarly reduced importin-α3 mRNA expression levels as upon TNF-α treatment (Figure 2N). It is important to note that IL-6 transcription is directly dependent on NF-κB, unlike IL-10, IFN-γ or TGF-β. These findings support the concept that TNF-α-activated NF-κB is crucial for importin-α3 gene expression. However, other cytokines might also reduce importin-α3 expression at high, but not low, cytokine concentrations (Figure S2C). Our findings support the concept that importin-α3 gene expression is controlled by TNF-α-activated NF-κB in a cell- and dose-dependent manner. This is in line with general control mechanisms in cells, such as negative or positive feedback loops, where gene transcription is regulated (e.g., either by activating or blocking the promoter, respectively, depending on demand) (Figure 2O). However, other, not yet defined, NF-κB independent pathways might also affect importin-α3 gene expression. Future studies are required to dissect the importin-α3 promoter for transcription factor binding sites other than NF-κB.

High-Level TNF-α-Inducing Human-Type HPAIV Infections Reduce Importin-α3 Protein Expression Levels in Human Respiratory Cells

Since the NF-κB transporting nuclear import factor importin-α3 was the only isofrom in the lung that altered its expression levels upon TNF-α treatment, we wanted to know how importin-α3 is regulated under respiratory stress associated with cellular cytokine changes. Therefore, we used an influenza infection model. We infected MEFs with highly pathogenic avian influenza A virus (H5N1) mutants that were known to act as low- or high-level TNF-α inducers (Figure 3A). Then, we infected MEFs either with the low-level TNF-α-inducing HPAIV containing an avian-like polymerase signature (SC35-PB2701D), common in influenza A virus (IAV) having a low replicative fitness in mammals, or its high-level TNF-α-inducing HPAIV counterpart containing a human-type polymerase signature (SC35M-PB2701N), common in IAV possessing a high replicative fitness and virulence.

Figure 2. Molecular Function and Regulation of Importin-α3

(A) Schematic overview of importin-α3-mediated NF-κB p65 nuclear translocation.
(B) Pull-down of NF-κB from lysates of untreated or TNF-α-treated A549 cells with Sepharose-immobilized GST-importin-α3 and α3. Expression of GST-importin-α3 isoform was verified with Coomassie staining (upper panel), and bound NF-κB p105, p65, and p50 were detected by western blot (lower panel). GAPDH protein detection served as loading control. One representative experiment is shown out of three independent experiments.
(C) Effect of TNF-α treatment on expression and localization of NF-κB p65 and importin-α3 in WT MEFs (n = 3).
(D and E) Nuclear fraction (NF; control p84) and cytoplasmic fraction (CF; control GAPDH) localization of NF-κB α3 in WT MEFs (n = 3). Reducing importin-α3 protein expression in NEMO−/−cells treated with TNF-α as compared to untreated cells (Figure 2M). Reduced importin-α3 expression in NEMO−/−cells could be restored upon transfection of a NEMO-expressing plasmid back into NEMO−/−cells (Figure 2M). Control treatment of NEMO−/−cells with IL-6, IL-10, interferon (IFN)-γ or TGF-β showed similarly reduced importin-α3 mRNA expression levels as upon TNF-α treatment (Figure 2N). It is important to note that IL-6 transcription is directly dependent on NF-κB, unlike IL-10, IFN-γ or TGF-β. These findings support the concept that TNF-α-activated NF-κB is crucial for importin-α3 gene expression. However, other cytokines might also reduce importin-α3 expression at high, but not low, cytokine concentrations (Figure S2C). Our findings support the concept that importin-α3 gene expression is controlled by TNF-α-activated NF-κB in a cell- and dose-dependent manner. This is in line with general control mechanisms in cells, such as negative or positive feedback loops, where gene transcription is regulated (e.g., either by activating or blocking the promoter, respectively, depending on demand) (Figure 2O). However, other, not yet defined, NF-κB independent pathways might also affect importin-α3 gene expression. Future studies are required to dissect the importin-α3 promoter for transcription factor binding sites other than NF-κB.
in mammalian species. Importin-α3 protein levels were strongly reduced in human-type HPAIV compared to avian-type HPAIV-infected MEfS (Figures 3B–3E). In contrast, protein expression levels of other importin-α isoforms (α1, α4, α5, and α7) were not affected (Figures 3F and 3G). Next, we wanted to assess whether replication of avian- and human-type HPAIV is affected in cells deficient in importin-α3. Cells deficient in importin-α3 were obtained from fibroblasts of mice generated for this study with a deleted importin-α3 gene (α3−/−; α3−/− MEfS; Figures S2D and S2E). MEfS with a deficient importin-α3 gene were obtained from mice with a deleted importin-α7 gene (α7−/−), described previously (Gabriel et al., 2011). Replication kinetics in MEfS revealed that avian-type but not human-type HPAIV replication is impaired in cells with a deleted importin-α3 gene (α3−/−), unlike in control cells, with a deleted importin-α7 gene (α7−/−) that acts as a positive factor of human-type HPAIV replication in mammalian cells (Hudjetz and Gabriel, 2012) (Figures 3H and 3I). Interestingly, avian-type HPAIV replication in α3−/− MEfS is increased by >1,000 times compared to WT control cells. Human-type SC35M-PB2701N HPAIV replication was comparable between WT and α3−/− cells (Figures 3J and 3K). In line, human-type H7N7 HPAIV replication was not affected in human H1299 α3−/− cells upon TNF-α treatment, likely due to the strong inhibition of importin-α3 protein expression (Figure 3L). In human H1299 α3−/− cells, SC35M-PB2701N HPAIV replication slightly increased, suggesting some residual inhibitory activity of importin-α3 in human cells (Figure 3L). These findings show that low-level TNF-α-inducing HPAIVs with avian-type polymerase signatures are highly sensitive to importin-α3-mediated virus restriction. In contrast, high-level TNF-α-inducing HPAIVs with human-type polymerase genes are resistant to importin-α3-mediated virus restriction.

**Importin-α3 Belongs to the Most Conserved Nuclear Transport Factors between Avian and Mammalian Species**

Differential regulation of importin-α3 expression levels upon avian- and human-type HPAIV infections prompted us to analyze the sequence identity of importin-α isoforms in various avian and mammalian species. Among all importin-α isoforms analyzed, importin-α1 was the most diverse nuclear import factor, with amino acid homology of 83.52% between chicken and humans (Figure 4A; Table S1). The isoforms importin-α3 and importin-α5 displayed the highest amino acid homologies between chicken and humans of 99.04% and 98.08%, respectively (Figures 4B and 4C; Table S1). Importin-α5, -α6, and -α7 presented amino acid homologies between chicken and humans of 95.91%, 94.22%, and 94.4%, respectively (Figures 4D–4F; Table S1). These findings suggest that particularly importin-α3 and importin-α4, both known nuclear transporters of NF-κB (Fagerlund et al., 2005), are highly conserved across species kingdoms. However, importin-α3 is expressed up to 5-times higher levels than importin-α4 in the murine and human lung (Figures 1C and 1D), highlighting particularly the importance of importin-α3 in respiratory immune responses. Thus, acquiring the ability to downregulate importin-α3 in the lung might serve as an important basis in avian-mammalian virus transmission. In order to see whether differential importin-α3 regulation is also present in human isolates, we analyzed human-type HPAIV strains that caused poultry outbreaks with human infections in the Netherlands (H7N7) or Thailand (H5N1). Infection of primary lung cultures with cytokine storm, including TNF-α-inducing H5N1 and H7N7 HPAIV isolates with human-type polymerase signatures (PB2627E; PB2701N), showed significantly reduced importin-α3 expression levels compared to their counterparts with avian-type polymerase signatures (PB2627E; PB2701D) (Figures 4G and 4H). These findings highlight the crucial role of importin-α3 as a respiratory factor conserved across animal kingdoms in HPAIV interspecies transmission.

**Human-Type Polymerase Signatures in HPAIV Mediate Reduced Importin-α3 mRNA Expression Levels in the Murine BE**

Next, we wanted to assess whether differential regulation of importin-α3 can also be detected in a murine infection model. Therefore, we assessed importin-α3 expression levels in lung samples of HPAIV-infected WT mice. Infection with avian-type SC35-PB2701D led to increased importin-α3 mRNA levels in the lung compared to uninfected controls on day 3 post infection (p.i.) (Figures 5A and 5B). In contrast, infection with human-type SC35M-PB2701N resulted in significantly reduced importin-α3 mRNA levels (Figure 5C). Since the human-type SC35M HPAIV contains mainly two host adaptive signatures (PB2 701N and NP 319K), we analyzed recombinant viruses with these single mutations to dissect their individual impact on the dysregulation of importin-α mRNA transcription. Herein, the PB2 D701N host adaptive mutation was predominantly responsible for reduced importin-α3 transcription, while its effect was further enhanced when combined with the NP N319K host

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**Figure 3. HPAIV Replication Kinetics in Importin-α3−/− Cells**

(A) TNF-α induction in MEfS infected with avian-type SC35-PB2701D or human-type SC35M-PB2701N H7N7 HPAIV (MOI = 0.1) measured by ELISA (n = 3). (B–E) Protein extracts thereof were analyzed and quantified by western blot at 48 h p.i. Importin-α3 and viral NP protein were detected (B and D) and quantified (C and E) with GAPDH (B and C) or NP adjustment (D and E) (n = 4). Dashed lines in (D) depict cropping of the relevant bands run on the same gel. (F and G) Additionally, importin-α1, -α4, -α5, and -α7 and viral NP protein were detected and quantified with GAPDH adjustment to confirm that observed effects were specific for importin-α3. Control-treated, uninfected samples or SC35-PB2701D-infected samples were set to 100%. Cross-reactivity of importin-α antibodies with NP resulted in a band (f) in (B) and (F). (H–K) Growth kinetics of avian-type SC35-PB2701D (H and I) and human-type SC35M-PB2701N (J and K) H7N7 HPAIV in importin-α−/− MEfS (n = 3). WT (black), importin-α3−/− (α3−/−, red), and importin-α7−/− (α7−/−, green) MEfS were infected with low (H and J) or high (I and K) multiplicities of infection (MOI). Virus titers given in plaque-forming units (p.f.u.) were determined by plaque assay on MDCKII cells (0, 24, 48, 72, and 96 h p.i.). (L) H1299 α3−/− (black) and α3−/− (red) cells were infected with human-type SC35M-PB2701N HPAIV (MOI = 0.001) and replication kinetics measured in the presence or absence of TNF-α (n = 3). Virus titers were determined by plaque assay on MDCKII cells (0, 24, 48 h p.i.). Data shown represent means of relative importin-α3 amounts or logarithmic virus titers ± SD of at least three independent biological experiments (p < 0.05; ***p < 0.001).
adaptive signature (Figures 5D–5F). Compared to all other importin-α isoforms, importin-α3 continuously showed the most prominent alterations. Interestingly, the most significant alterations in importin-α3 mRNA levels were detected at 3 days p.i. compared to 1 day p.i. (Figures S2F–S2K), consistent with the concept that accumulating cytokine levels through multiple replication kinetics is required for a profound cytokine-dependent repression of importin-α3 transcription. Avian-type SC35-PB2701D virus replication, which caused an increase in importin-α3 mRNA levels, was restricted to the respiratory tract (Figures S3A and S3B). In contrast, human-type SC35M-PB2701N viral replication led to a significant reduction of importin-α3 mRNA levels in the lung, increased virus load in the lung, and systemic virus spread (Figure S3C). The recombinant viruses with the single or combined human-type PB2 701N and NP 319K signatures showed stepwise systemic spread along with their ability to reduce importin-α3 mRNA levels in the lung (Figures S3D–S3F). These findings further highlight that host
adaptive signatures mediate impaired importin-α3 transcription, resulting in high-level virus replication.

To determine the respiratory cell types, where reduction of importin-α3 expression occurs, we stained serial lung sections for importin-α3 as well as influenza virus antigen. Importin-α3 was strongly expressed in the BE of avian-type SC35-PB2701D-infected mice, correlating with reduced virus-antigen-positive cells in these tissues compared to human-type SC35-M-PB2701N-infected animals (Figures 5G and 5H). In contrast, importin-α3 protein levels in SC35-M-PB2701N-infected mice were predominantly reduced in the BE, which correlated with high-level viral replication (Figures 5G and 5H). In line, virus replication was increased in the lungs of α3−/− mice compared to their α3+/+ WT littermates infected either with SC35-PB2701D or SC5M-PB2701N virus (Figure S4). Thus, differential regulation of importin-α3 protein expression predominantly occurs in the BE, the key site of productive influenza virus replication (Abdel-Ghafar et al., 2008; Peiris et al., 2007).

Failure to Induce NF-κB-Controlled Antiviral Gene Expression in the Lung Leads to Increased Mortality in Importin-α3-Deficient Mice

Next, we addressed the question of whether reduction of pulmonary importin-α3 levels would affect NF-κB-mediated antiviral gene expression and whether this would affect disease outcome. Therefore, we infected α3−/− mice and their WT littersmates (α3+/+) with avian- or human-type HPAIV. Lung transcriptomics of mice infected with human-type SC35-M-PB2701N virus.
Figure 6. Genes Responsive to HPAIV Infection in Lungs of WT and Importin-α3−/− Mice

WT and importin-α3−/− (α3−/−) mice were infected with SC35-PB2701D (H7N7) or SC35M-PB2701N (H7N7). Control mice received PBS.

(A) Heatmap depicting the row Z scores of the normalized expression values of 143 differentially expressed genes (p value ≤ 0.01 and |log2FC| ≥ 1.5) that were clustered according to their expression profiles in lungs on day 3 p.i.

(B) Heatmap depicting row Z scores of the normalized expression values of 79 genes with a previously described role in innate immunity (Engels et al., 2017).

(C) TNF-α expression levels detected by transcriptome analysis on day 3 p.i. in lungs of infected α3+/+ mice relative to the maximum expression observed. Control was set to 100%.

(D) TNF-α protein levels measured by ELISA on day 1 p.i. in α3+/+ and α3−/− mice. Data show average results obtained from pooled lung samples of three WT mice per virus strain (*p < 0.05; ***p < 0.001).

(E and F) Relative mRNA-expression levels of antiviral genes under an NF-κB promoter determined by qRT-PCR 3 days p.i. in control (E) and infected lungs (F) of α3+/+ and α3−/− mice. Relative expression values of α3+/+ for each gene normalized to HPRT were set to 1. Data shown represent mean ± SD (n = 5; **p < 0.01; ***p < 0.001).

(G and H) Survival in α3+/+ and α3−/− mice infected with avian- (G) or human-type (H) H7N7 HPAIV. Data shown represent means ± SD (n = 9 for SC35 and n = 4 for SC53M).
HPAIV in WT as well as in α3−/− mice revealed a subset of substantially altered expression patterns in genes involved in innate immunity (Engels et al., 2017) (Figure 6A; Tables S2 and S3). Herein, genes controlled by an NF-κB promoter were particularly prominent, with 34% of all analyzed genes (27 out of 79 hits) (Figure 6B). Moreover, literature analysis of the hits highlighted that especially genes involved in TNF pathways were of high abundance, supporting the in vitro findings (data not shown). TNF-α mRNA levels were increased in the lungs of human-type HPAIV-infected α3−/− mice when compared to lungs of avian-type HPAIV-infected animals (Figure 6C). As expected, TNF-α protein expression, which is also regulated by NF-κB, was reduced in α3−/− mice infected with both avian- and human-type HPAIV (Figure 6D). We then assessed a subset of antiviral genes under an NF-κB promoter that showed reduced expression in α3−/− mice, such as TNFSF-10, IFN-β1, IFN-β2, MX-1, IFN-γ, EIF2AK2 (alias PKR), IFR-3, MAVS, and IFNA-14. While none of these genes showed significant alterations in PBS-treated control mice irrespective of the presence or absence of the importin-α3 gene (Figure 6E), the transcription, most prominently of IRF-2 and MX-1 genes involved in major antiviral pathways, was most significantly reduced in SC35M-PB2701N HPAIV-infected α3−/− mice compared to α3+/+ mice (Figure 6F). Infection of α3−/− mice with both SC35-PB2701N and SC35M-PB2701N resulted in enhanced weight loss and lethality compared to α3+/+ mice (Figures 6G, 6H, and 5S). Virulence was significantly increased in human-type HPAIV-infected α3−/− mice (Table S4). These data show that importin-α3 is required for the efficient expression of antiviral genes under the control of an NF-κB promoter. Failure to induce these antiviral pathways leads to high HPAIV virulence in mice.

**DISCUSSION**

Importin-α proteins serve as key proteins that regulate nucleocytoplasmic cargo protein transport to maintain cellular homeostasis. Disruption of balanced transport pathways, caused under stress or inflammatory conditions, can cause serious diseases. In immortalized cancer cell lines, importin-α1 expression levels are particularly high compared to non-cancerous cells, as shown here as well as in many other studies before. Here, we shed light on the anatomic distribution of importin-α isoforms in the mammalian lung. We provide evidence of the highly abundant expression of importin-α3 in the murine and human RT. Moreover, importin-α isoforms are expressed as a gradient, with increasing concentrations from the URT to the LRT. Abundant and exclusive high-level expression of importin-α3 was a hallmark of the lung, unlike in other solid organs assessed, such as the liver, spleen, and brain. Knowledge on the importin-α expression pattern in the lung is important to understand the molecular basis of cellular homeostasis that might be dominated by importin-α3-transported transcription factors.

According to the crystal structure of importin-α3, which was recently solved, it was postulated that it is highly selective of cargo proteins, unlike other importin-α isoforms (Pumroy et al., 2015). The high substrate specificity is reflected by only a few cargo proteins reported to be transported into the nucleus by importin-α3. Importin-α3 seems to exclusively serve as a transporter of p53, a transcription factor involved in many cancer pathways (Marchenko et al., 2010). Importin-α3 is also a key factor in the transport of the NF-κB subunits p50 and p65 into the nucleus along with importin-α4, which is the closest relative of importin-α3 (Fagerlund et al., 2005). Mutations in ARM3 of the importin-α3 protein were shown to be crucial for p50 binding, while mutations in ARM7 and 8 were shown to be required for p65 binding (Fagerlund et al., 2005). Based on the crystal structure of importin-α3, it was further postulated that it might pose a “fast track” nuclear import pathway under certain circumstances, such as environmental stress (e.g., viral infections), where cellular amounts of importin-β1 may be limited, requiring fast counteraction (Pumroy et al., 2015). This hypothesis is in line with our findings in this study suggesting that importin-α3 serves as a defense mechanism against influenza by promoting the nuclear entry of NF-κB, which controls the transcription of many antiviral genes.

We further show that importin-α3 gene expression is highly dynamic under inflammatory conditions. We identified an NF-κB binding site within the importin-α3 promoter element and show that its transcription is dependent on TNF-α-activated NF-κB in a concentration-dependent manner. HPAIVs that induce low-level TNF-α expression, such as those with avian-type polymerase signatures (PB2 627K and 701D), increase importin-α3 mRNA expression presumably by NF-κB induction. In contrast, high-level TNF-α expression inducing HPAIV, such as those harboring human-type polymerase signatures (PB2 627K and 701N), significantly reduces importin-α3 mRNA expression levels by a yet-unknown mechanism. One possibility would be that high numbers of NF-κB proteins may block the promoter of the importin-α3 gene and thereby its transcription. Other possible interferences could be with NF-κB-activating complexes, such as NEMO or its repressor, IκBα. Future studies are required to unravel the detailed mechanism underlying the failure of importin-α3 promoter activation. Such diverse feedback loops are known in cellular processes, where gene transcription depends on demand. Herein, generating a bottleneck in importin-α3 availability leads to failure in inducing an adequate antiviral immune response, as shown in mice lacking the importin-α3 gene. Interestingly, more than 30% of the differentially expressed genes between α3−/− mice and their α3+/+ litter mates were under the control of an NF-κB promoter, highlighting the significant contribution of importin-α3 in NF-κB-mediated gene transcription. We could further show, using HPAIV strains of the H5N1 and H7N7 subtypes that have caused pneumonia and death in humans, that downregulation of importin-α3 mRNA transcription was a hallmark that was mediated by their human-type polymerase signatures. The introduction of avian-type polymerase signatures into the genetic backbone of these HPAIV outbreak strains restored importin-α3 mRNA transcription in the lung. It is important to note that these human-type HPAIVs induce a general cytokine storm, including TNF-α. Thus, more detailed studies are required to understand the impact of other cytokines on the transcription level of importin-α3. However, a pulmonary increase in TNF-α levels with a concomitant increase in virulence has been described for many human H5N1 and H7N7 isolates containing human-type (PB2627K) signatures (Abdel-Ghafar et al., 2008; de Jong et al.,...
2006; Hatta et al., 2001; Peiris et al., 2007; Szretter et al., 2007). Based on these findings, we propose that human-type HPAIVs have evolved mechanisms within the mammalian host to evade antiviral response pathways by high-level induction of TNF-α-activated NF-κB, which blocks the transcription of importin-α3, the major classical nuclear import protein expressed in the mammalian lung.

We have shown previously that influenza A viruses utilize differential importin-α isoforms to ensure replicative fitness (Gabriel et al., 2011). Upon zoonotic transmission of HPAIV from birds to humans, they switch from importin-α3 to importin-α7 usage. While importin-α3 acts as an inhibitor of influenza virus polymerase, importin-α7 acts as a positive factor of influenza virus replication. Importin-α7 is believed to act in a transport-independent function by a yet-unknown mechanism. Until now, it was unclear how importin-α3 mediates antiviral activity. To unravel the mechanism underlying the importin-α3-mediated inhibition, the generation of cell culture models as well as of a murine model with a deleted importin-α3 gene was essential. Our previous observation that replication of HPAIVs with avian-type polymerases, unlike HPAIV with human-type polymerases, was restricted in cells silenced using importin-α3 siRNA and is likely due to residual importin-α3 expression sufficient to induce nuclear import of NF-κB and thus the transcription of various antiviral genes. Furthermore, our findings here support the concept that avian influenza viruses evade importin-α3 restriction by blocking its transcription. As a consequence, influenza viruses get access to the second-most abundant isoform in the lung, importin-α7, to ensure high-replicative fitness in the human lung (Gabriel et al., 2011).

Finally, we here provide key insights on the dynamic role of the respiratory importin-α3 protein as an “immune sensor” of viral infections in the lung. Attempts to restore depleted importin-α3 levels in the lung might pose new strategies to treat severe respiratory virus infections.

**STAR METHODS**

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**SUPPLEMENTAL INFORMATION**

Supplemental Information can be found online at https://doi.org/10.1016/j.celrep.2020.107549.

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**AUTHOR CONTRIBUTIONS**


**DECLARATION OF INTERESTS**

The authors declare no conflicts of interest.


**STAR METHODS**

**KEY RESOURCES TABLE**

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**LEAD CONTACT AND MATERIALS AVAILABILITY**

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Guelsah Gabriel (guelsah.gabriel@leibniz-hpi.de). Materials, reagents and mouse lines of this study are available with a completed Material Transfer Agreement.
EXPERIMENTAL MODEL AND SUBJECT DETAILS

Human Subjects
Biopsy material from human uninfected URT (nasal concha) and LRT (lung, mainly consisting of alveolar tissue) was obtained from the Erasmus Medical Center in Rotterdam, the Netherlands, from donors at the Erasmus Medical Center in Rotterdam. Approval to use the clinical samples for scientific use was obtained from the Dutch Medical Ethical Committee under the permit numbers MEC-2011-129, MEC 2008-207 and MEC 2009-128. Donor sex and age is unknown.

Cells and Viruses
Cell lines of human embryonic kidney cells (HEK293T, ATCC), human alveolar adenocarcinoma cells (A549, H1299, ATCC), African green monkey kidney cells (VeroE6) and human cervical adenocarcinoma cells (HeLa, ATCC) were grown in Dulbecco’s modified Eagle’s medium (DMEM, PAA) supplemented with 10% fetal calf serum (FCS; PAA), 1% penicillin/streptomycin (P/S, PAA), and 1% L-Glutamine (PAA). Mouse embryonic fibroblasts (MEF, see description of their generation in a separate paragraph below) were grown in DMEM supplemented with 10% FCS, 1% P/S, 1% L-Glutamine, 1% sodium pyruvate, and 1% non-essential amino acids (NEAA, PAA). Immortalized NEMO+/− MEFs (Makris et al., 2000) were kindly provided by Michael Karin (University of California, San Diego). Madin Darby canine kidney cells (MDCKII, ATCC) were grown in Minimal Essential Medium (MEM, PAA) supplemented with 10% FCS, 1% P/S, and 1% L-Glutamine. Primary human bronchial (HBEpC, PromoCell, C-12640) and primary human small airway epithelial cells (hSAEpC, PromoCell, C-12642) were grown in Airway Epithelial Cell Media or Small Airway Epithelial Cell Media (PromoCell), respectively, supplemented with 1% P/S. HBEpC cell donor was a 52 year old female subject. HSAEpC cell donor was a 57 year old female subject. Cells were cultivated at 5% CO₂, 96% rH and 37°C (PromoCell), respectively, supplemented with 1% P/S. HBEpC cell donor was a 52 year old female subject. Cells were cultivated at 5% CO₂, 96% rH and 37°C. Recombinant H7N7 HPAIV viruses SC35-PB2701N, SC35M-PB2701N, the single-point mutants SC35-PB2701N and SC35-NP319K or the double mutant SC35-PB2701N-NP319K (Gabriel et al., 2011) as well as recombinant A/Thailand/1(KAN-1)/2004 (H5N1) wild-type strain (H5N1-PB2701N) (Gabriel et al., 2011; Puthavathana et al., 2005), the avian-type single point mutant thereof (H5N1-PB2701D) and H7N7 HPAIV virus isolates A/Netherlands/219/2003 (H7N7-PB2627K, fatal case) or the recombinant virus thereof (H7N7-PB2627E) (Czudzai-Matwich et al., 2014; Foucheir et al., 2004) were used.

All experiments with H5N1 and H7N7 viruses were conducted at the Heinrich Pette Institute, Leibniz Institute for Experimental Virology in Hamburg, Germany, in biosafety level 3 facilities, except for the SC35M-PB2701N strain that was classified as biosafety level 2 agent, and approved by the relevant German authorities (Gentechnikbehörde Hamburg).

Animals
Importin-α3−/− mice in the C57BL/6 genetic background and WT littermates thereof were bred and housed under specific pathogen-free conditions at the animal facility of the Heinrich Pette Institute, Leibniz Institute for Experimental Virology, Hamburg, Germany. We used 6-12 week old female and male mice. Experimental groups were matched regarding equal numbers of female and male mice as well matched in comparable age groups.

Animal experiments were performed in strict accordance with the guidelines of the German Animal Welfare Regulations. All animal protocols were approved by the relevant German authority (Behörde für Stadtentwicklung und Umwelt Hamburg, license numbers: 29/09, 54/11). Mice were humanely killed upon > 25% weight loss or upon achievement of defined body index scores (e.g., on physical appearance, clinical signs, behavior, eating and drinking etc.) according to the German Animal Welfare Regulations.

METHOD DETAILS

Transfection and Vectors
Transfections were performed using Lipofectamine 2000 (Invitrogen) or jetPRIME® (Polyplus-Transfection) according to the manufacturer’s instructions. Vector constructs used were pHW2000-(SC35-PB2701D, SC35-PB2701K, SC35-PA, SC35-NP, SC35M-PB2701N, SC35M-PB1, SC35-MA, SC35M-NP, SC35M-PB2701N, SC35-MA, SC35M-NP) (Gabriel et al., 2005), pcDNA-importin-α1-N-FLAG, pcDNA-importin-α3-N-FLAG, pcDNA-importin-α25-N-FLAG and pcDNA-importin-α7-N-FLAG, pPol-I-NP-Luc-human (Gabriel et al., 2005), pRL-TK (Promega), pLightSwitch_Prom-control and pLightSwitch_Prom-importin-α3 (renilla luciferase reporter constructs; Switchgear Genetics), pFL-TK and pCMVTAG-NEMO (Addgene). The pcDNA-importin-α3-N-FLAG constructs were described previously (Gabriel et al., 2011; Wang et al., 1997). pFL-TK was generated by amplification of the firefly luciferase gene from the pPol-I-NP-luc-human construct (Gabriel et al., 2005) using primers p5_H领衔_Fluc_for (5′-GGCGTGACCATGGAGACGCACAAAAACATAAAGAA AGGCCGCCGC-3′) and p3_Fluc_NotI_rev (5′-GGCGCGCCGCTTTACCTTGGTACCTTGGCATTC-3′). Insertion of the firefly luciferase gene into pRL-TK without the renilla luciferase gene was performed via the Nhel and Notl restriction sites.

Antibodies
Primary antibodies used for western blot and immunohistochemical analyses include goat anti-importin-α1 (Abcam, ab6036), rabbit anti-importin-α1 (Abcam, ab54489), goat anti-importin-α3 (Abcam, ab6039), rabbit anti-importin-α4 (Abcam, ab6038), rabbit anti-importin-α3 (Novus Biologicals, #NB100-93345, replaced by #NB100-81651), rabbit anti-importin-α3 (Köhler et al., 1997), rabbit anti-importin-α5/α7 (Köhler et al., 1999), rabbit anti-NF-κB p65 (Cell Signaling, #3012), rabbit anti-LSD1 (Cell Signaling, #2139), rabbit
anti-GAPDH (Cell Signaling, #2118), mouse anti-nuclear matrix protein p84 (Abcam, ab487), mouse anti-FLAG M2 (Sigma-Aldrich, #F3165), rabbit anti-HA (Sigma-Aldrich, #H908), mouse anti-influenza A virus nucleoprotein (HB65; American Type Culture Collection, Manassas, VA) antibodies and rabbit anti-FPV serum (Gabriel et al., 2008). Anti-rabbit-HRP, anti-goat-HRP, anti-mouse-HRP (Sigma-Aldrich, #A8275, #A5420, and #A4416, respectively), and IRDye-conjugated anti-rabbit (Li-Cor) antibodies were used as secondary antibodies for western blot.

**Importin-α sequence analysis**

Nucleotide sequences of the coding regions of importin-α1, importin-α3, importin-α4, importin-α5, importin-α6 and importin-α7 were downloaded from the NCBI nucleotide database for *Homo sapiens* (GenBank: NM_001320611.1, NM_002268.5, NM_002267.4, CR456743.1, NM_001366307.2, NM_012316.5), Macaca fascicularis (GenBank: NM_001283904.1, XM_005546256.2, XM_005585866.2, XM_005547987.2, XM_005551660.2, XM_005544119.2), Sus scrofa (GenBank: NM_001163404.1, XM_021069735.1, NM_001193575.1, NM_001163405.1, XM_005564393.3, XM_021095806.1), *Mus musculus* (GenBank: NM_010655.3, NM_008464.4, NM_008466.5, BC006771.1, NM_008468.4), Gallus gallus (GenBank: NM_001006209.1, NM_001007963.1, NM_001193575.1, NM_001030774.1, NM_015284586.2, NM_001012841.2) and Anas platyrhynchos (GenBank: XM_005012314.3, XM_027464723.1, XM_027471420.1, XM_027449713.1, XM_027454663.1). For *Mus musculus*, no importin-α6 sequence was available, while the importin-α7 sequence was missing for *Anas platyrhynchos*. The nucleotide sequences were subsequently translated into amino acid sequences using the standard genetic code. For each Importin-α, the sequences were aligned using Muscle version 3.8.1551 (Edgar, 2004) with standard parameters. Pairwise nucleotide and amino acid sequence identities were then calculated using Clustal Omega version 1.2.4 (Sievers et al., 2011).

**Generation of Importin-α3/C0 Mice and Importin-α/C0 MEFS**

To generate an importin-α3 targeting construct, a 900 bp-long sequence upstream and a 5100 bp-long sequence downstream of exon 1 of the importin-α3 gene were cloned into a targeting vector described before (Walther et al., 1998). After homologous recombination in embryonic stem (ES) cells, exon 1 bearing the translational start site for the importin-α3 protein, was deleted. ES cells were electroporated with the linearized construct and clones were picked after double selection with neomycin and gancyclovir. Positive clones were identified by PCR and one of them was chosen for blastocyst injection. From the injected cell clone, germine chimeras were obtained and bred with C57BL/6 mice. mutant ES cell clones, germline chimeras were obtained and bred with C57BL/6 mice. Importin-α3 deficient mice were backcrossed for eight generations to the C57BL/6 genetic background; the colony was maintained by breeding heterozygous mice. For genotyping, the following primers were used for PCR on genomic DNA of tail or ear biopsies: Impa3for 5'-GCCCACTCATAGCGGAATG-3'; Impa3rev 5'-GACAACGAGAAATTGGACAACC-3'. To confirm the gene deletion, absence of importin-α3 mRNA and protein was determined by RT-PCR and western blot of organs. Absence of importin-α3 mRNA with β-Actin as reference was verified using the following primers: Impa3Ex3_for 5'-GCCCACTCATAGCGGAATG-3'; Impa3Ex3_rev 5'-GACAACGAGAAATTGGACAACC-3'; Impa3Ex3_for 5'-GACAACGAGAAATTGGACAACC-3'; Impa3Ex3_rev 5'-GACAACGAGAAATTGGACAACC-3'.

**Generation of Importin-α3/C0 Cell Lines**

To generate H1299 importin-α3/C0- and control importin-α3/C0+ cell lines, the cells were seeded in 12 wells and transfected using Lipofectamine 2000 (Invitrogen) with either a control CRISPR-Cas9 plasmid or an importin-α3/C0 CRISPR-Cas9 plasmid (Santa Cruz Biotechnology). 48 hours post transfection positive cell clones were sorted via FACS into 96 well plates with help of a green fluorescent protein selection marker. Clones were tested via western blot for importin-α3 expression and positive clones were cultivated further for cryopreservation.

**Animal Experiments**

Mice were anaesthetized with ketamine-xylazine (70 mg/kg and 7 mg/kg, respectively) and intranasally infected with the respective virus dose in 50 µl PBS. WT mice were infected with 6×10⁵ p.f.u. (100-fold MLD₉₀ of SC35M-PB₂7₀₁N in BALB/c mice) of SC35-PB₂7₀₁N. SC35-PB₂7₀₁N, the single-point mutants SC35-PB₂₇₀₁N or SC35-NP₁₉₁K or the double mutant SC35-PB₂₇₀₁N-NP₁₉₁K. Control groups received PBS. On day 3 p.i., five animals per time point were anaesthetized with an overdose of ketamine-xylazine (140 mg/kg and 14 mg/kg, respectively) and sacrificed after intracardial perfusion with DEPC-treated 1x PBS. Perfused organs (lung, trachea, liver, spleen, brain) were removed and dissected into pieces (< 5 mm) and submerged in RNA later RNA Stabilization Reagent (QIAGEN). Murine URT samples were obtained by decapitation of the head after the anterior end which includes the nasal concha was excised and submerged in RNA later. For relative mRNA-Expression levels determination of antiviral genes under an NF-κB promoter by RT-qPCR, WT (n = 5) and importin-α3/C0- (n = 5) mice were infected with 10⁶ p.f.u. of SC35M. Control groups received PBS. On day 3 p.i., five animals were anaesthetized with an overdose of ketamine-xylazine and sacrifice. Whole lungs were removed and dissected into pieces and submerged...
in RINaVer. Organs (lung, spleen, brain) of uninfected WT mice (n = 3) for western blot lysate preparation were obtained after short inhalative isoflurane narcosis followed by cervical dislocation. After dissection into small pieces, the organs were immediately frozen on dry ice.

For virus titer determination of organs, WT (n = 3) and importin-α3/-/- (n = 3) mice were infected with 6 10^6 p.f.u. of SC35-PB2701D (~0.1x MLD50 of SC35-PB2701D in C57BL/6 mice) or a low and high dose of SC35M-PB2701N (10^5 p.f.u.; ~0.5x MLD50 or 6*10^4 p.f.u.; ~30x MLD50 of SC35M-PB2701N, respectively). On days 3 and 6 p.i., three animals per time point were sacrificed, organs (lung, trachea, brain) removed, virus titers determined by plaque assay 64, and the URT and LRT stained immunohistochemically against influenza virus and importin-α antigen. For this, murine formalin-fixed, paraffin-embedded (FFPE) lungs or skinned, decalcified FFPE heads were thin sectioned (3 μm). Heads were decalcified in 10% EDTA (pH 7,4) for 8 days, followed by sagittal division and incubation in same solution for 3 more days. Mouse lethal dose 50 (MLD50) was assessed by infecting WT (n = 4-9) and importin-α3/-/- (n = 4-9) mice with serial virus dilutions of SC35-PB2701D (10^2, 5*10^2, and 10^3 p.f.u.) or SC35M-PB2701N (10^2, 10^3, and 10^4 p.f.u.). Animals were monitored for weight and survival for 14 days p.i. and the MLD50 was calculated. Humane endpoints were defined according to the German Animal Welfare Regulations as described above.

**Immunohistochemical Analysis (IHC-P)**

Viral and importin-α, -α3, and -α7 antigen were determined by IHC-P with antigen-specific primary and HRP-conjugated secondary antibodies. URT and lungs of infected WT and importin-α3/-/- mice were excised on days 1 and 3 p.i.. FFPE thin-sections (3 μm) of murine samples and human biopsies were prepared as described (Gabriel et al., 2009). Viral antigen was stained using the mouse anti-influenza A nucleoprotein (HB65; American Type Culture Collection, Manassas, VA) or the rabbit anti-FPV-serum, a biotin-conjugated anti-rabbit secondary antibody (Jackson ImmunoResearch. #711-066-152), the Avidin/Biotin Blocking Kit SP2001 (Biozol), and the ZytoChemPlus (HRP) Broad Spectrum (DAB) Kit (Zytomed) according to the manufacturer’s instructions. For visualization of importin-α isoforms in murine and human FFPE samples, the described rabbit anti-importin-α, rabbit anti-importin-α3 (Novus Biologicals), and rabbit anti-importin-α5/α7 antibodies were used together with a biotin-conjugated anti-rabbit secondary antibody (DAKO; #E0432), a peroxidase-conjugated avidin-biotin-complex-reaction kit (DAKO), and the HRP substrate AEC (DAKO). Counter-staining was performed with hematoxylin. Since the importin-α7 antibody, cross-reacts with importin-α5, lung tissues of mice with a deletion of either the importin-α5 or the α7 gene were additionally used as controls (data not shown). To exclude false-positive staining due to cross-reactivity of the antibodies, FFPE thin sections were additionally stained using importin-α isofrom-specific blocking peptides for importin-α1 (aa518-529, QVQDGAPGFNF, GenScript), importin-α3 (aa509-521, NSSANVPTEGFOF, Abcam, #ab23144), importin-α5 (aa3-16, TPGKENFRLKSYKN, GenScript) and importin-α7 (aa3-12, MASPGKDNYR; aa526-536, PEAP-MEGFQL, GenScript) in different concentrations (0.2 μg/ml, 2 μg/ml, and 20 μg/ml) (Table S5). Primary antibodies were incubated with the respective peptides for 30 min prior to IHC-P. No unspecific staining was observed with 2 μg/ml and 20 μg/ml of peptide (data not shown). Images were acquired using a NIKON AZ100 wide field light microscope with a NIKON Digital Sight DS-Ri1 high-resolution microscope camera, AZ Plan Fluor 5x and AZ Plan Apo 1x objectives, and the NIKON Digital Sight DS-L2 Imaging Controller. Image processing was performed using Adobe® Photoshop® CS4 Extended v.11.0.2.

**Laser Microbeam Microdissection (LMM)**

LMM was performed on murine lung cryosections to obtain bronchiolar epithelium and alveolar tissue samples using the PALM MicroBeam System (Axiovert 200 microscope incl. PALM CapMover, PALM RoboStage 1, PALM RoboSoftware v2.2.; P.A.L.M. Microlaser Technologies GmbH/ZEISS). After thoracotomy of WT C57BL/6 mice, lungs were filled with Tissue Tek® Compound (50% in 1x PBS, Sakura Finetek) and were shock-frozen in melting isopentane. Cryosections (10 μm) were generated at the Cryostat CM3050 S (LEICA), let dry for 5 min at ~20°C and stained with cresyl violet (2 min 70% ethanol, ~20°C; 2 s 1% cresyl violet in 50% ethanol, RT; 30 s 70% ethanol, ~20°C; 30 s 100% ethanol).

**RNA Isolation**

Total RNA from cells or tissue samples was isolated using the innuprep RNA Minikit (Analytik Jena). Homogenization of ~50 mg of RNAwere-fixed organs was performed in 450 μl lysis buffer RL with 10 sterile, stainless steel beads (Ø 2 mm, #22.455.0010, Retsch) at 30 Hz and 4°C for 6 min in the mixer mill MM400 (Retsch). RNase free DNase Set (QIAGEN) was used for on-column DNase l-treatment. After elution in RNase-free water, 1 U/μl Ribolock RNase Inhibitor (Thermo Scientific) was added. RNA concentration and purity were determined using the Nanodrop 1000 (Peqlab). Microdissected murine tissue was vortexed for 1 min in lysis buffer RL, incubated for 30 min at RT and vortexed again (1 min). After short centrifugation, the lysate was stored at ~80°C overnight. After thawing, 1 min of vortexing, and centrifuging (8 s), the sample was mixed ten times with 700 μl of 70% ethanol using a needle and syringe. RNA was isolated according to the innuprep RNA Minikit’s instructions. Total RNA from human FFPE URT was isolated from 1-2 sections (20 μM) after xylene deparaffinization using the QIAGEN RNeasy® FFPE Kit (QIAGEN). Isolated RNA was either processed immediately or stored at ~80°C.

**cDNA Generation**

cDNA from tissues or cells was generated using random primers and the SuperScript III Reverse Transcriptase (Invitrogen) according to the manufacturer’s instructions. Random nonamer primer (Gene Link, pd(N)9, 26-4000-06, final concentration: 5 μM) and dNTP
cDNA from microdissected samples was generated using the RevertAid H Minus Reverse Transcriptase (Thermo Scientific). 2 μl of total RNA were treated with 1 U/μl Dnase I and 1x Dnase I buffer (Sigma-Aldrich, #AMPD1) for 15 min at RT (reaction volume: 10 μl). DNA restriction was stopped by adding 1.5 μl 50 mM EDTA and incubation for 10 min at 70°C. Then, 1x Reaction Buffer, DTT (final concentration: 500 μM), dNTP-Mix (Promega; final concentration: 500 μM each dNTP), 250 ng random hexamer primer (pd(N)6, Eurofins MWG Operon), and 100 U RevertAid H Minus Reverse Transcriptase were added (final reaction volume: 20 μl). cDNA was generated using the cycle 42°C for 50 min, 70°C for 15 min, and 4°C for ∞.

For cDNA from human FFPE URT, 250 ng of total RNA were incubated with Random Primers (Promega, 500 μg/ml, # C1181; final concentration: 500 ng) and dNTP mix (final concentration: 500 μM each dNTP) for 5 min at 65°C in a volume of 13 μl. After cooling (at least 1 min on ice), 1x First Strand Buffer, DTT (final concentration: 5 mM), RNasin® Ribonuclease Inhibitor (Promega, #N251B; final concentration: 2 U/μl) and 10 U/μl SuperScript III Reverse Transcriptase were added (final reaction volume: 20 μl). cDNA was generated using the following cycle: 25°C for 5 min, 50°C for 45 min, 70°C for 15 min, and 4°C for ∞. All cDNA samples were processed immediately or stored at −20°C.

Design of RT-qPCR Primers
DNA oligonucleotides (Sigma-Aldrich; Table S6 for RT-qPCR were designed using Clone Manager 9 Professional Edition or Primer-BLAST (https://www.ncbi.nlm.nih.gov/tools/primer-blast/index.cgi).

Real-Time Quantitative PCR (RT-qPCR)
Gene of interest (GOI) mRNA expression levels were determined using specific primer pairs (Table S6). Singleplex reactions (10 μl) were set up manually in UltraPure DNase/RNase-Free Distilled Water (GIBCO) in MicroAmp® Optical 96-Well Reaction Plates (Invitrogen, #4306737): 5 μl Platinum® SYBR® Green qPCR SuperMix-UDG (2x, Invitrogen), 0.02 μl ROX Reference Dye (25 μM, Invitrogen, final concentration: 50 nM), 300 nM of forward and reverse primer each, and 1 μl cDNA template. RT-qPCR runs were conducted on the ABI 7500 Fast System (Applied Biosystems) in the Standard 7500 mode with endpoint fluorescence detection: 2 min at 50°C, 3 min at 95°C, 50 amplification cycles (15 s at 95°C, 10 s at 65°C, and 30 s at 72°C). Analysis was performed in triplicate or quadruplicate for each GOI and GAPDH in each sample. Subsequently, melting curve analysis was performed on the ABI 7500 Fast System (15 s at 95°C, 1 min at 60°C, 15 s at 95°C). Correct amplicon size was checked by agarose gel electrophoresis. Data of reactions with false products was excluded from data analyses. Relative expression values were determined using the E−ΔΔCT-method (Ramakers et al., 2003; Ruijter et al., 2009). The Rn-values were exported from the SDS Software v1.3.1 (Applied Biosystems) to Microsoft Office Excel 2007 and N0-values for the starting concentration of the transcript in the original sample were obtained with LinReg PCR Software v1.1.1 (Ruijter et al., 2009). The averaged N0-value of the GOI (e.g., N0 (importin-α2b), n = 3–4 technical replicates) was then normalized with the averaged N0-value for GAPDH (N0 (GAPDH)) of the respective sample. The relative N0 GOI/N0 (GAPDH)-expression values of the biological replicates are presented. For easier comparison of differential expression, the averaged relative N0-value was set 1. Relative mRNA expression levels of NF-κB regulated genes were determined using specific primer pairs (Table S6) for the genes of interest (GOIs) and the reference gene HPRT. Singleplex reactions (20 μl) were set up manually in H2O PCR grade (ROCHE®) in LightCycler® 480 Multiwell Plate 96 (ROCHE®, #04729692001): 10 μl FastStart Essential DNA Green Master (2x, ROCHE®), 300 nM of forward and reverse primer each, and 2 μl cDNA template. RT-qPCR runs were conducted on the LightCycler® 96 Real-Time PCR System (ROCHE®) with endpoint fluorescence detection: 10 min at 95°C, 50 amplification cycles (15 s at 95°C, 10 s at 65°C, and 20 s at 72°C). Analysis was performed in triplicate for each GOI and HPRT in each sample. Subsequently, melting curve analysis was performed (15 s at 95°C, 15 s at 60°C, 1 s at 95°C). Correct amplicon size was checked by agarose gel electrophoresis. The relative N0 GOI/N0 (HPRT)-expression values of the biological replicates are presented.

Protein Extraction and Purification
Whole cell lysates (WCL) of cells were obtained by lysis in HEPES lysis buffer (50 mM HEPES (pH 8.0), 200 mM NaCl, 0.5% IGEPAL, 25% glycerol, 1 mM PMSF, 0.07 μl/ml β-mercaptoethanol, 1x HALT Protease and Phosphatase Inhibitor Cocktail, 1x EDTA Solution) or samples were fractionated into cytoplasmic (CF) and nuclear fractions (NF) using the NE-PER Nuclear and Cytoplasmic Extraction Reagents (Thermo Fisher Scientific Inc.). Whole organ lysates of uninfected WT mice were obtained by weighing ~30 mg of tissue into O-ring tubes filled with 8 stainless steel beads (Ø 2 mm, #22.455.0010, Retsch). After washing the organ pieces once with 800 μl 1x PBS and centrifugation at 500 xg for 45°C for 5 min, the supernatant was discarded and 10 μl HEPES lysis buffer per mg organ were added. Homogenization was carried out at 4°C and 20 Hz for 2 min followed by 30 Hz for 1 min in the mixer mill MM400 (Retsch). The homogenates were then incubated for 5 min on ice and subsequently centrifuged for 20 min at 21,000 xg and 4°C. The supernatant was transferred into a fresh tube, mixed with 4x protein loading dye and boiled for 5 min at 95°C before being subjected to western blot analysis. Whole cell lysates of HEK293T cells transfected with 10 μg of each of the pcDNA-importin-α-N-FLAG constructs were subjected to affinity purification using the EZview Red ANTI-FLAG M2 affinity gel (Sigma-Aldrich) and eluted using a 3x FLAG peptide.
(Sigma-Aldrich) according to the manufacturer’s instructions. Thus, purified FLAG-tagged importin-α isoforms were obtained and used to generate standard curves to determine the affinities of the different isoform-specific importin-α antibodies. The protein amounts of each isoform were quantified in murine organ homogenates using western blot analyses and standardized using the individual standard curves as described previously (Köhler et al., 2002, 1999).

Western Blot
Successful knockdown of importin-α isoforms was confirmed by western blot analysis using the described goat anti-importin-α1, goat anti-importin-α3, goat anti-importin-α4, and rabbit anti-importin-α5/α7 antibodies, respectively, with normalization to GAPDH using the rabbit anti-GAPDH antibody. The same antibodies were used for detection and quantification of endogenous importin-α isoforms in cell or organ lysates. For detection and quantification of FLAG-tagged importin-α isoforms, the mouse anti-FLAG M2 antibody was used. Amounts of viral NP protein or NF-κB p65 in cell lysates after viral infection or TNF-α treatment were quantified using the rabbit anti-FPV serum and the rabbit anti-NF-κB p65 antibodies, respectively. For detection of HA-tagged NEMO, the rabbit anti-HA antibody was used. Equal amount of protein was loaded for each sample using GAPDH, p84 or LSD1 as loading control. Quantification of protein was performed with the Bioimager Image Quant LAS 4000 at non-saturated levels with GAPDH or LSD1 adjustment for WCL, GAPDH adjustment for CF, and LSD1 of p84 adjustment for NF. Alternatively, NP levels were adjusted and used for normalization.

Immunofluorescence
WT MEFs as well as MEFs deficient for importin-α3 were seeded in glass bottom dishes (#81218-200, Ibidi GmbH) using regular growth medium (10% FBS, 1% P/S, 1% L-Glutamine, 1% NEAA, 1% sodium pyruvate). After 8h, the growth medium was replaced with starvation medium (serum reduced to 0.5% FBS) and cells were starved for another 16h. Subsequently, cells were either control-treated or treated with murine TNF (10 ng/ml; #575204, BioLegend, Inc) for 15 min at 37°C. Then, immunofluorescence staining was performed. All washing steps were performed with phosphate buffered saline (PBS), and all incubation steps were performed at room temperature, if not otherwise stated. Briefly, cells were washed and fixed with 4% paraformaldehyde (in PBS) for 10 min at 37°C. Fixed cells were washed twice and permeabilized with 0.1% Triton X-100 (in PBS) for 20 min. After washing, cells were blocked with 3% bovine serum albumin (BSA) solution (in PBS) and then incubated with a primary antibody recognizing murine NF-κB p65 subunit (1:300 in 3% BSA/PBS; #8242S, Cell Signaling) for 1h. Cells were washed again and incubated for 1h with a donkey anti-rabbit IgG antibody coupled to Alexa-555 dye (1:500 in 3% BSA/PBS; A31572, Life Technologies GmbH). This step also included staining of nuclei with Hoechst dye (1:1000 in 3% BSA/PBS; #62249, Life Technologies GmbH). After further washing, stained cells were stored in PBS at 4°C. Images were acquired on a confocal Nikon Eclipse Ti-E spinning disc microscope (Nikon, Japan) at 100-fold magnification. Image processing was performed using Nikon NIS-Elements Advanced Research (version 4.51; Nikon, Japan) and ImageJ software (National Institute of Health, NIH). Semi-automatic quantification of nuclear and cytoplasmic NF-κB p65 signals was carried out using a custom-designed plugin for the ImageJ software which is available on request for the plugin. Data are based on three independent biological replicates, each performed with technical duplicates.

HPAIV Growth Kinetics
WT MEFs and MEFs with deleted importin-α3 or -α7 genes were infected with different multiplicities of infection (MOI) of SC35-PB2701D (H7N7, MOI = 0.1 or 0.001) or SC35M-PB2701N virus (H7N7, MOI = 0.01 or 0.0001). 7*10^5 cells were inoculated with 2 mL virus dilution a WT MEFs and MEFs with deleted importin-

Library Preparation and Next Generation Sequencing
Lungs of WT and α3−/− mice infected with 6*10^7 p.f.u. of SC35-PB2701D or SC35M-PB2701N (H7N7) were subjected to gene expression analysis. Control mice received PBS. On day 3 p.i., whole lungs were perfused, harvested and submerged in RNAlater and total RNA was isolated using the innuPrep RNA Mini Kit (Analytik Jena). RNA samples from three animals per mouse strain and virus were pooled and used for gene expression analysis. For each sample, 1μg of total RNA was used for generating sequencing libraries with the Illumina TruSeq RNA Sample Preparation Kit v2 as recommended by the manufacturer (Illumina Inc.). Size and quality of
the libraries were visualized on a BioAnalyzer High Sensitivity DNA Chip (High Sensitivity DNA Kit, Agilent Technologies). Diluted libraries (2 nM) were multiplex-sequenced on the Illumina HiSeq 2500 instrument. For each sample between 78 and 113M, paired-end reads of 101 bp length were generated. The reads were aligned to the murine reference transcriptome (UCSC mm10) using Bowtie2 (v2.2.2) (Langmead and Salzberg, 2012). DESeq (Anders and Huber, 2010) was employed to assess differential expression based on read counts per gene. Genes were hierarchically clustered according to their expression profiles using the R function hclust (http://www.R-project.org/). DAVID (v6.7) (Huang et al., 2009) was employed to functionally annotate the resulting clusters. Networks of protein-protein interactions were computed and visualized using Cytoscape (Shannon et al., 2003).

TNF-α ELISA

TNF-α protein amount was determined in lung homogenate supernatants of WT mice infected intranasally with 6×10^4 p.f.u. of SC35-PB2701D or SC35M-PB2701N viruses (H7N7). Whole lungs were harvested on day 1 p.i. and ~50–100 mg of tissue homogenized with ~300 μl glass beads (RETSCH #22.222.0003) in 1 mL 1x PBS (PAA) in the mixer mill MM400 (Retsch; 20 Hz, 4°C, 10 min). After centrifugation (6000 xg, 4°C, 5 min), lung homogenate supernatants were stored at -80°C. Supernatants of three animals per virus were pooled. Then, the Enzyme-linked Immunosorbent Assay (ELISA) Kit for murine TNF-α (USCN Life Science Inc.) was used to determine TNF-α concentrations.

TNF-α Treatment, Importin-α3 mRNA Expression, Subcellular Localization and Promoter Activation

HSAEpC, WT MEFs and NEMO–/– MEFs were seeded in 24-well plates until confluent or in 10 cm-dishes for 24 h, respectively, and starved for 24 h. Then, cells were control-treated (w/o) or treated with human TNF-α (50 ng/ml; #210-TA, R&D Systems) for 6 h or with murine TNF-α (50 ng/ml; #575204, BioLegend, Inc.) for 6 h or 24 h, respectively. WCL, CF, and NF were subjected to western blot analysis for importin-α3 and NF-κB p65 detection. Total RNA were isolated 6 h post treatment for relative importin-α3 expression, using RT-qPCR.

For determination of the relative importin-α3 mRNA expression levels after cytokine treatment, NEMO–/– MEFs were seeded in 10 cm cell culture dishes (5×10^6 cells per dish). After serum starvation for 24 h, cells were control-treated (mock) or treated 10 min with 10ng/ml of murine TNF-α, IL-6, IL-10, IFN-γ and TGF-β. After 6 h post treatment, cells were harvested, RNA isolated and cDNA generated before RT-qPCR analysis.

For the determination of nuclear import of NF-κB p65 after TNF-α stimulation, WT MEF and Importin-α3 KO MEF were seeded in 10 cm cell culture dishes (1×10^6 cells per dish). After serum starvation for 18 h, cells were treated with 10ng/ml TNF-α. Subsequently, cells were harvested by scraping in ice-cold PBS, centrifuged and after removing the supernatant, the cells were resuspended in 50μl of Buffer A (10mM Tris pH7.9; 1.5mM MgCl₂; 10mM KCl). After adding 6μl of 2% NP-40 and vortexing, the samples were incubated for 10 min on ice and centrifuged for 15 min at 6,000rpm at 4°C. The supernatant (cytosolic fraction) was placed in a fresh tube, subsequently the pellet was washed again with 100μl of buffer A, centrifuged again for 15 min and the supernatant was removed. For nuclear fractions, the pellet was resuspended in 50 μl of buffer B (20 mM Tris pH 7.9; 10mM KCl; 1.5mM MgCl₂; 20% Glycerol) and vortexed; then 33 μl of buffer C (20mM Tris pH 7.9; 1.2M KCl; 1.5 mM MgCl₂; 20% Glycerol) was added and the samples were incubated on a wheel at 4°C for 45 min. Afterward, samples were centrifuged for 30 min at 12,000 rpm at 4°C and the supernatant (nuclear fraction) was transferred to a fresh reaction tube. Both fractions were mixed with 4x protein loading dye and boiled for 5 min at 95°C before being subjected to western blot analysis.

For determination of importin-α3 promoter activation in MEF cells, cells were seeded in 96-well plates (5000 cells/well) to reach approximately 60% confluency after 24h. Then, cells were transfected using FuGene HD transfection reagent (SwitchGear Genomics), either with pLightSwitch-Prom-Control (SwitchGear Genomics; 100 ng/well) as a control, or pLightSwitch-Prom-importin-α3 (SwitchGear Genomics; 100 ng/well) for importin-α3 promoter activity readout. After 24h, transfected cells were treated with murine TNF-α, IL-10 or TGF-β at 0, 1 or 50 ng/ml, for 15 min. Promoter activity was measured at 3h post treatment using LightSwitch™ Luciferase Assay Reagent (SwitchGear Genomics, LS010) according to manufacturer’s instructions.

For determination of importin-α3 promoter activation, A549 or HSAEpC were seeded in 96-well plates for 24 h or until 80% confluency, respectively, and transfected with pLightSwitch-Prom-Control (50 ng; Control) or pLightSwitch_Prom-importin-α3 (50 ng; α3) and pFL-TK (250 ng; firefly luciferase transfection control) using Lipofectamine-2000 or jetPRIME®, respectively. 14 h after transfection, serum starvation was initiated for 30 h. Subsequently, cells were treated with 0, 10, 50 or 100 ng/ml (A549) or with 0, 1, 10 or 100 ng/ml (HSAEpC) of human TNF-α and lysed 20 h later. Firefly and Renilla luciferase activities were assessed using the Dual-Luciferase® Reporter Assay System (Promega, E1910).

NEMO Complementation Assay

NEMO–/– MEFs were transfected in suspension with 5 μg empty vector pcDNA3.1 or pcDNA-HA-human NEMO in 6-well plates using Lipofectamine 2000 (Invitrogen) for 6 h. At 48 h post transfection, cells were serum starved for 24 h. Then, cells were control-treated (w/o) or treated with murine TNF-α (50 ng/ml; #575204, BioLegend, Inc.) for 6 h. WCL were subjected to western blot analysis for importin-α3 and NEMO detection.
**QUANTIFICATION AND STATISTICAL ANALYSIS**

P values of RT-qPCR data were obtained by Kruskal-Wallis one-way ANOVA using PAST v.2.17c (http://folk.uio.no/ohammer/past/). RT-qPCR data were considered as relevant when more than ±1.5-fold change was detected in relative mRNA expression levels. Statistical significance of animal survival rates was determined with GraphPad Prism 5 v.5.03 (Graphpad Software, Inc.) using the Log Rank (Mantel-Cox) test and the Kaplan-Meier survival curves. All other mean values, standard deviations (SD), and P values were calculated with GraphPad Prism 5 (v5.03) or the GraphPad Software, Inc. tool (https://www.graphpad.com/quickcalcs/ttest1/?Format=SD) using the unpaired, two-tailed Student’s t test or the one-sample, two-tailed Student’s t test when all values in one group were identical, e.g., set 100%. Statistical significance was defined as p < 0.05 (* p < 0.05, ** p < 0.01, *** p < 0.001).

**DATA AND CODE AVAILABILITY**

Transcriptome sequence data reported in this publication is submitted to the European Nucleotide Archive (ENA) and is available under following link: https://www.ebi.ac.uk/ena/data/view/PRJEB8023.