Title
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Permalink
https://escholarship.org/uc/item/5hc01844

Journal
PLoS pathogens, 15(8)

ISSN
1553-7366

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Publication Date
2019-08-05

DOI
10.1371/journal.ppat.1007460

Peer reviewed
The NDV-3A vaccine protects mice from multidrug resistant Candida auris infection

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Abstract

Candida auris is an emerging, multi-drug resistant, health care-associated fungal pathogen. Its predominant prevalence in hospitals and nursing homes indicates its ability to adhere to and colonize the skin, or persist in an environment outside the host—a trait unique from other Candida species. Besides being associated globally with life-threatening disseminated infections, C. auris also poses significant clinical challenges due to its ability to adhere to polymeric surfaces and form highly drug-resistant biofilms. Here, we performed bioinformatic studies to identify the presence of adhesin proteins in C. auris, with sequence as well as 3-D structural homologies to the major adhesin/invasin of C. albicans, Als3. Anti-Als3p antibodies generated by vaccinating mice with NDV-3A (a vaccine based on the N-terminus of Als3 protein formulated with alum) recognized C. auris in vitro, blocked its ability to form biofilms and enhanced macrophage-mediated killing of the fungus. Furthermore, NDV-3A vaccination induced significant levels of C. auris cross-reactive humoral and cellular immune responses, and protected immunosuppressed mice from lethal C. auris disseminated infection, compared to the control alum-vaccinated mice. The mechanism of protection is attributed to anti-Als3p antibodies and CD4+ T helper cells activating tissue macrophages. Finally, NDV-3A potentiated the protective efficacy of the antifungal drug micafungin, against C. auris candidemia. Identification of Als3-like adhesins in C. auris makes it a target for immunotherapeutic strategies using NDV-3A, a vaccine with known efficacy against other Candida species and safety as well as efficacy in clinical trials. Considering that C. auris can be resistant to almost all classes of antifungal drugs, such an approach has profound clinical relevance.

Author summary

Candida auris has emerged as a major health concern to hospitalized patients and nursing home subjects. C. auris strains display multidrug resistance to current antifungal therapy
following competing interests: JE, and AI are founders and shareholders of NovaDigm Therapeutics. All other co-authors have no formal association with NovaDigm. The findings and conclusions in this manuscript are those of the authors and do not necessarily represent the official position of the Centers for Disease Control and Prevention or NIH.

and cause lethal infections. We have determined that C. auris harbors homologs of C. albicans Als cell surface proteins. The C. albicans NDV-3A vaccine, harboring the N-terminus of Als3p formulated with alum, generates cross-reactive antibodies against C. auris clinical isolates and protects neutropenic mice from hematogenously disseminated C. auris infection. Importantly, the NDV-3A vaccine displays an additive protective effect in neutropenic mice when combined with micafungin. Due to its proven safety and efficacy in humans against C. albicans infection, our studies support the expedited testing of the NDV-3A vaccine against C. auris in future clinical trials.

Introduction
The fungus Candida auris was first detected in 2009 from an ear canal infection in Japan [1]. However, the earliest known strain of C. auris dates back to 1996 isolated in a retrospective analysis of previously misdiagnosed samples from Korea [2]. Since then, C. auris has been reported in more than 20 countries, with a significant number of cases detected in the Unites States [3]. Patients can remain colonized with C. auris for long periods and the yeast can persist on surfaces in healthcare environments, which results in spread of the organism between patients in healthcare facilities [4, 5]. Clinical isolates of C. auris have been recovered from a variety of specimen types, including normally sterile body fluids, wounds, mucocutaneous surfaces, and skin [4]. However, bloodstream infection remains the most commonly observed clinical manifestation of C. auris, with alarming in-hospital global crude mortality rates of 30 to 60% [6, 7]. Further and of high importance, some isolates of C. auris exhibit multidrug resistance with elevated MICs to all three major antifungal classes, including azoles, echinocandins, and polyenes, resulting in limited treatment options [8]. The ability of C. auris to persist and survive in an environment outside the host is unique from most other pathogenic Candida species. This characteristic of survival in a hostile environment perhaps means that C. auris has virulence determinants that help it adapt, adhere, and persist in those settings. Recent reports reveal that C. auris can adhere to polymeric surfaces and form highly drug resistant biofilms [5, 9, 10]. In fact, similar to other Candida infections, presence of a central venous catheter has been identified as a risk factor for C. auris [4, 11].

We embarked on a study to determine if C. auris possessed evolutionarily conserved adhesin protein homologs similar to those found in another phylogenetically-related human fungal pathogen C. albicans. Using bioinformatic and structural homology modeling, we discovered that C. auris contained protein homologs of the C. albicans Als3p. Als3p, a member of the agglutinin-like sequence (Als) family of proteins, is also known to be conserved in other non-albicans Candida species [12–14], and has a multifunctional adhesin and invasin properties essential for host pathogenesis [15, 16]. The N-terminus of C. albicans Als3p has been developed as a vaccine that induces protective antibody and cell-mediated immune responses [17, 18], and has been shown to be safe and efficacious in a clinical trial against recurrent vulvovaginal candidiasis [17, 19–24]. We found that NDV-3A vaccination induced C. auris cross-reactive antibody and T cell responses. Moreover, the sera from NDV-3A vaccinated mice blocked virulence characteristics of C. auris and enhanced opsonophagocytic killing of this fungus. Furthermore, vaccination with NDV-3A antigen protected mice from lethal C. auris infection and improved the efficacy of sub-therapeutic doses of micafungin. The afforded protection appears to be mediated by humoral and cellular immunity that leads to fungal clearance by tissue macrophages. These findings identify NDV-3A as a promising vaccine for adjunctive treatment of life-threatening bloodstream infections caused by the multidrug resistant C. auris.
Results

C. auris possesses homologs of C. albicans Als3

Bioinformatic and 3-D structural modeling strategies identified three conserved proteins in C. auris with homologies to C. albicans Als3p. In particular, these C. auris Als3p homologs, PIS50650.1, PIS50263.1 and XP_018167572.2, displayed ~30% identity and ~50% similarity (with several regions of homology reaching up to 88%) to C. albicans Als3p. Furthermore, sequence alignments revealed that all three proteins possessed domains characteristic of Als3p, such as an N-terminal secretory signal sequence, central amyloid forming and repeated Ser/Thr-rich sequences, as well as the presence of a C-terminus GPI anchor. (Table 1, S1 Dataset). Once identified, the above sequences were assembled to produce 3-D structural models for analysis in relation to Als3p (Swiss-model figures shown in Fig 1). Based on this modeling strategy, C. albicans Als3p shares striking similarity to the C. auris proteins, particularly in the N-terminus motif (Fig 1, S1 Dataset).

Anti-Als3p anti-sera from NDV-3A-vaccinated mice recognize C. auris

Because bioinformatic analysis revealed considerable sequence as well as 3-D structural homology between C. albicans Als3p and homologous proteins of C. auris, and predicted cell surface localization of these homologs, we hypothesized that anti-Als3p antibodies should recognize C. auris in vitro. The anti-Als3p antibodies were generated by vaccination of mice with NDV-3A, a vaccine based on the N-terminus of C. albicans Als3p, which is known to induce high titers of serum anti-Als3p antibodies [18, 20, 24]. Sera from these alum-vaccinated mice were examined in two different binding assays in vitro. First, different C. auris clinical isolates (CAU-01, 03, 05, 07 and 09) and C. albicans cells were grown for 90 min under germ-tube inducing conditions since Als3p is expressed on C. albicans hyphae [25], and then treated with anti-NDV-3A sera, followed by immunostaining with fluorescent labeled secondary IgG. Immunostaining of germinating C. albicans cells (positive control), confirmed the specificity of Als3p to the C. albicans filaments. C. auris yeast cells do not differentiate into hyphae, yet anti-Als3p antibodies bound to the cell surface of the fungus, as depicted by a diffused or punctate green fluorescence of the yeast cells. Fungal cells treated with sera from alum-vaccinated mice failed to fluoresce (Fig 2 panel A). The binding of anti-Als3p antibodies to five strains of C. auris obtained from different clades showed the universal presence of Als3p homologs in this multidrug resistant yeast (See Table B in S1 Table for drug susceptibility). Fungal cells treated with sera from alum-vaccinated mice failed to fluoresce (Fig 2 panel A).

Next, the extent of anti-Als3p antibody binding to the fungal cells was quantified by flow cytometry. The anti-Als3p antibodies present in the NDV-3A-vaccinated sera bound significantly to C. auris isolates (indicated by a shift in the peak representing mean fluorescence intensity), while the negative control did not. These results demonstrate the specificity of anti-Als3p antibodies to C. auris and further validate the modeling strategy which previously revealed the homology of C. albicans Als3p to Als-like proteins on C. auris (Fig 2 panel B [upper histograms]).

To determine which of the three Als3p homologous protein cross reacted to ant-Als3p antibodies, we produced gene deletion mutants of C. auris PIS50650.1, PIS50263.1 and XP_018167572.2 using CRISPR (S1 and S2 Figs). All three individual Als3p homolog deletion mutants of CAU-09 (C. auris) showed reduced binding to anti-Als3p IgG antibodies compared to wild type CAU-09 strain, indicating their role in cross-reactivity to NDV-3A vaccine (Fig 2 panel B [lower histograms]).
NDV-3A vaccine against Candida auris

Table 1. List of the Candida albicans [SC5314] Als3p orthologs in Candida auris with percent homology and structural features.

| S. No | NCBI Protein Database Sequence ID | Name | Length | Match | Range | Identities | Positives | GPI-Anchor Location | Ser/Thr Repeats | Amyloid Sequences |
|-------|----------------------------------|------|--------|-------|-------|------------|-----------|-------------------|----------------|-----------------|
| 1     | PIS50650.1                        | Hypothetical protein CJD97_004175 [Candida auris] | 1804 | 1     | 1 to 371 | 124/381 (33%) | 187/381 (49%) | C-23 | Yes | Yes |
| 2     | PIS50263.1                        | Hypothetical protein CJD97_004954 [Candida auris] | 835 | 1     | 2 to 348 | 111/356 (31%) | 179/356 (50%) | C-17 | Yes | Yes |
| 3     | XP_018167572.2                    | Hypothetical protein QG37_06265 [Candida auris] | 861 | 1     | 8 to 431 | 115/435 (26%) | 204/435 (46%) | C-21 | Yes | Yes |

1*Protein amino acid homology was analyzed by NCBI BLAST.
2GPI-anchor location was predicted by GPI-SOM tool (http://gpi.unibe.ch/).
3Serine or Threonine rich repeats were detected by using RADAR tool (https://www.ebi.ac.uk/Tools/pfa/radar/).
4Amyloid sequences were detected by PASTA 2.0 tool (http://protein.bio.unipd.it/pasta2/).

NDV-3A vaccination induced robust C. auris cross-reactive antibodies and T-cell responses

We have previously shown that NDV-3A vaccine mediated protection against C. albicans infection required both Als3p-specific antibodies and CD4+ Th1/Th17 immune responses, and anti-Als3p antibody titer threshold predicts protective efficacy [21]. Thus, we reasoned...
that robust antibody and T cells responses would be critical for protection against *C. auris* in *vivo*. Therefore, we examined the magnitude of anti-Als3p antibodies binding to *C. auris* anti-gens by using cell-based ELISA. First, we confirmed that NDV-3A-vaccinated sera harbored high levels of anti-Als3p antibodies, by using ELISA coated with recombinant Al3p N-terminus. As expected, NDV-3A-vaccinated mice sera contained high titers of anti-Als3p antibodies (mean end-point titer of 12,800), while alum vaccinated sera had none. Next, we developed a
cell-based ELISA of *C. albicans* or *C. auris* to compare the magnitude of anti-Als3p antibodies to antigens of yeast cells. We allowed the binding of serially diluted mouse sera to *C. albicans* germ tube cells or *C. auris* yeast cells and detected the endpoint titer with HRP-conjugated anti-mouse IgG detection antibodies. We found that NDV-3A vaccinated mice had high titers of cross-reacting antibodies against *C. albicans* cells, with titers similar to the recombinant Als3p N-terminus-based ELISA (12,800). Consistent with the confocal microscopy and flow cytometry results, and compared to sera obtained from alum-vaccinated mice, NDV-3A-vaccinated mice sera were found to have cross-reactive anti-Als3p antibodies to *C. auris* cells albeit with 4-fold less than *C. albicans* (mean end-point titer of 3,200) (Fig 3 panel A).

We also examined whether NDV-3A vaccination induced memory CD4+ T cells that are also cross-reactive to *C. auris*. Splenocytes from NDV-3A- or alum-vaccinated mice were stimulated with either recombinant Als3p-N-terminus or heat-killed *C. auris* for 5 days, followed by treatment with PMA/ionomycin in the presence of monesin/brefeldin A protein transport inhibitor. These cells were stained with fluorescent-labelled antibodies against surface markers and intracellular cytokines. Flow cytometry analysis of these stained cells revealed a significant presence of high frequency of Als3p-specific and *C. auris* cross-reactive CD4+ T cells, which were comprised of Th1 (IFN-γ +), Th2 (IL-4+) and Th17 (IL-17+) cells (Fig 3 panel B, S3, S4, S5 and S6 Figs). The *C. auris* cross-reactive T cells were not detected in alum control mice. Further, the total *C. auris* or Als3p-specific CD4+ T cell responses were similar in magnitude and were slightly biased towards Th2 type. Finally, the *C. auris* -specific Th17 cell response was higher compared to Als3p-specific responses (Fig 3 panel B, S3, S4, S5 and S6 Figs). Overall, these results revealed that, NDV-3A induced robust *C. auris* cross-reactive antibodies as well as CD4+ T cell immune responses.

**Anti-NDV-3A sera reduced *C. auris* biofilm formation and enhanced opsonophagocytic killing of *C. auris* by macrophages**

Attachment of microorganisms to an abiotic surface is the first step in formation of a drug-resistant biofilm. Biofilm formation has been most studied in *C. albicans* and Als3p is essential for establishment of the early stages of biofilm growth [26]. Recently, we have reported that blocking Als3p by anti-Als3p antibodies abrogate *C. albicans* biofilm formation [20]. *C. auris* also has the capacity to form drug-resistant biofilms [9]. Thereby, we evaluated if anti-Als3p antibodies from NDV-3A-vaccinated sera could similarly inhibit biofilm formation in *C. auris*. Sera from NDV-3A-vaccinated mice significantly inhibited biofilm formation compared to control anti-alum sera (*p* = 0.001) (Fig 4 panel A). Of note, sera were used at 1:10 (sera to media) ratio, and heat-treated prior to their contact with fungal cells, to rule out the role of complement in the inhibitory activity (Fig 4 panel A).

We previously demonstrated that anti-Als3p antibodies act as an opsonin to enhance phagocyte-mediated killing of *C. albicans* [20]. Thus, we posited that Als3p antibodies would enhance sensitivity of *C. auris* to macrophage killing. To test this hypothesis, heat-inactivated mice sera (at 10% concentration) from NDV-3A- or alum-vaccinated mice were incubated with *C. auris* to allow binding of anti-Als3p antibodies to the fungal cell surface, and then added to mice macrophage cell-line (1:1). As a control, *C. auris* yeast cells were subjected to macrophages without any sera. Both NDV-3A- and alum-vaccinated mice sera did not have any negative impact on growth of *C. auris* (S7 Fig). However, sera from NDV-3A-vaccinated mice significantly enhanced macrophage-mediated killing of *C. auris* compared to sera from alum vaccinated mice (40% vs 2% killing respectively, normalized to *C. auris* killing by macrophage-only condition, *p* = 0.028). Thus, anti-Als3p antibodies generated by the NDV-3A vaccine enhanced the opsonophagocytic killing of *C. auris* (Fig 4 panel B).
NDV-3A vaccination protected mice from lethal *C. auris* disseminated infection

Blood stream infections are predominant manifestations of *C. auris*, accompanied by significant mortality in susceptible patients. Since, anti-Als3p antibodies generated by NDV-3A vaccination bound to *C. auris*, blocked virulence and rendered it susceptible to macrophages, we reckoned that NDV-3A vaccination would have the potential to protect against *C. auris* disseminated candidiasis. We developed a neutropenic mouse infection model by screening different clinical isolates of *C. auris* (CAU-01, 03, 05, 07, 09) for their lethality in mice. An intravenous 5x10^7 CFU/mouse dose of *C. auris* CAU-09 showed 100% mortality in neutropenic mice and was used for vaccine efficacy testing (immunocompetent mice are resistant to *C. auris* bloodstream infection). To test the vaccine-mediated protective efficacy, mice were either
vaccinated with NDV-3A or alum, three times (primary + two booster doses), and then infected with *C. auris* via the tail vein (Fig 5 panel A). Mice vaccinated with alum displayed 100% mortality by day 7, while NDV-3A-vaccinated mice showed 40% overall survival. Surviving mice appeared healthy at 21 days when the experiment was terminated (Fig 5 panel B). Furthermore, while both sets of vaccinated mice displayed weight loss post infection compared to uninfected mice, NDV-3A-vaccinated mice had lesser extent of weight loss versus alum-vaccinated mice (Fig 5 panel C). In replicate studies to the survival experiments, mice were sacrificed 4 days post infection, and their target organs (kidneys and brain) were collected for tissue fungal burden. The NDV-3A-vaccinated mice had a 10-fold lower (~1.0 log) fungal burden in both kidneys and brains compared to alum control mice (*p* = 0.0006) (Fig 5 panel D).

Histopathological examination of tissues collected from mice sacrificed on day 4 post infection, confirmed inhibition of disseminated fungal infection in kidneys of NDV-3A-vaccinated mice, while alum-treated mice had multiple abscesses containing *C. auris* cells throughout the organ (Fig 6 panel A). Compared to kidneys, the brain had an overall 100-fold reduced fungal burden (Fig 5 panel D), which was also apparent in the histopathological images (Fig 6). While overall only a few cells could be visualized in the brain, alum-vaccinated mice appeared to have more fungi compared to brains of NDV-3A-vaccinated mice. Interestingly, in both cases, *C. auris* cells were localized in the blood capillaries, rather than the brain tissue itself (Fig 6 panel B).

**NDV-3A vaccine mechanism of protection is antibody- and CD4+ T helper cell-dependent and requires tissue macrophages**

To dissect the role of NDV-3A vaccine-mediated protection, we conducted serum adoptive transfer and immune cell depletion experiments. Briefly, pooled anti-NDV-3A (mean titer = 25600, S8 Fig) or alum serum was administered i.p. to naïve neutropenic Balb/c mice.
Fig 5. NDV-3A-vaccinated mice are protected from *C. auris* infection. (A) Outbred CD-1 mice were immunized three times with either NDV-3A or alum alone. Neutropenia was induced by two doses of cyclophosphamide and cortisone acetate on day -2 and +3 of infection. After two weeks of final boost, the mice were infected through tail vein with lethal $5 \times 10^7$ CFU/mouse dose. The mice from both groups were divided into two arms. One arm was observed for survival for 21 days, and another arm were sacrificed on day 4 post infection. (B) NDV-3A vaccinated mice were protected from *C. auris* dissemination infection. The mice survival was compared by using Mantel-cox Log-rank Test ($n = 10$ mice/group). (C) After 4 days post infection, weight of NDV-3A or alum control vaccinated mice were compared ($n = 8$ mice/group). (D) On day 4 post infection, NDV-3A or alum...
starting on day -1 and weekly thereafter, relative to i.v. infection with *C. auris*. The mice that received anti-NDV-3A serum had significantly enhanced median survival time and 21-day % survival compared to the mice that received anti-alum serum (6 days and 33% survival for anti-NDV-3A vs. 4 days and 0% for anti-alum serum, p = 0.0006) (Fig 7 panel A). Since anti-NDV-3A antibodies enhanced the opsonophagocytic killing of the *C. auris* by murine macrophages *ex vivo*, we hypothesized that these antibodies help in controlling *C. auris* infection through macrophage-mediated killing in the neutropenic mouse model. To examine this possibility, we vaccinated mice as previously described and depleted their macrophages by injecting clodronate-liposome. Mice receiving PBS-liposome served as placebo control. Macrophage depletion in primary lymphoid organs was verified in a separate set of mice by staining of spleen and inguinal lymph node cells with macrophage-specific phenotypic markers (CD11b + F4/80+). By day 4 after the second injection of clodronate-liposome, the macrophage population decreased by ~97% in spleens taken from neutropenic mice administered clodronate-liposome when compared to PBS-liposome treat neutropenic mice, while lymph nodes maintained a very low frequency of macrophages in both depletion and control depletion groups (S9 Fig). Interestingly, macrophage depletion completely reversed the survival benefit afforded by the NDV-3A vaccine (from 37% survival to 0% survival, and from 11 days median survival to 5 median survival days, p value = 0.01). This clearly indicates that loss of macrophages in neutropenic mice severely compromised the NDV-3A vaccine-mediated protection against systemic *C. auris* infection (Fig 7 panel B).

Next, to examine the role of CD4+ helper T cells, we vaccinated mice with NDV-3A or alum as previously described and depleted CD4+ T cells by injecting anti-mouse CD4 (GK1.5 IgG2b) antibodies through i.p. route on day -3 and 0 relative to infection. The mice were made neutropenic on day -2 and infected with *C. auris* on day 0 followed by survival analysis for 3-weeks post-infection. The CD4+ T helper cell depletion was verified on day 4 after the second injection of anti-mouse CD4 antibodies in primary lymphoid organs as described previously [27]. Mice treated with anti-CD4 antibodies had a 99% reduction in the frequency of CD4+ T cells compared to isotype control antibody treated mice (S10 Fig). Further, CD4+ T cell depletion in NDV-3A vaccinated mice had marked reduction in median survival time against systemic *C. auris* challenge (12 vs 5.5 median survival days for CD4+ replete and CD4+ deplete arms [p = 0.04 by Mantel Cox comparisons] and 38% vs 13% survival, [p = 0.18 by Log Rank test]). Interestingly, the protection afforded by the NDV-3A vaccine in absence of CD4+ was still higher than survival of mice vaccinated with alum with or without depletion of CD4+ cells (13% vs 0% survival, and 6 vs 4 median survival days for NDV-3A vaccinated and CD4+ deplete vs. alum vaccinated mice, p value = 0.0028) (Fig 7 panel C).

Collectively, these adoptive and targeted depletion studies show both antibodies (humoral) and CD4 cells are required in providing protection against systemic *C. auris* infection in this neutropenic mouse model, likely via enhancing fungal-killing by tissue macrophages.

**NDV-3A vaccination potentiated the activity of micafungin, against *C. auris* disseminated infection**

In the clinical setting, any therapeutic vaccine approach is likely to be used in combination with standard antifungal therapy. Given that NDV-3A vaccination provided significant protection...
Fig 6. Comparison of histopathology of kidneys and brain tissue sections from NDV3-A or alum-vaccinated and C. auris infected mice. Tissue sections obtained on 4 days after the infection were stained with PAS. (A) Kidneys. The kidney from the control alum mouse had numerous abscesses with visible C. auris fungal cells throughout the tissue. (B) Brain. NDV-3A-vaccinated mouse did not show any visible abscesses or C. auris organism but alum mouse brain was visibly infected with C. auris. The arrow facing down show the C. auris in the tissues, and arrows facing up show the blood capillaries.

https://doi.org/10.1371/journal.ppat.1007460.g006
against *C. auris* candidemia, we questioned if vaccination could potentiate the activity of antifungal drugs, *in vivo*. To investigate this possibility, we vaccinated mice with NDV-3A or alum as above, made them neutropenic and then infected them with *C. auris*. Twenty-four hours post infection, one group of NDV-3A-vaccinated or alum control mice were treated with sub-inhibitory concentrations of micafungin (0.5 mg/kg body since CAU-9 is micafungin sensitive [MIC = 0.125 μg/ml]) daily for 1 week. As expected, NDV-3A protected 40% of mice from succumbing to infection, while micafungin provided only 20% protection. Interestingly, when used in combination, NDV-3A and micafungin significantly enhanced the overall survival of mice to >70% (p = 0.04, compared to NDV-3A-vaccination alone; and p = 0.001, compared to micafungin alone). These results show that NDV-3A vaccine acts additively with active antifungals in protecting mice from *C. auris* candidemia (Fig 8).
Discussion

The simultaneous global emergence of the pathogenic yeast *C. auris*, its ability to persist in healthcare-settings, combined with its potential to resist all classes of antifungal drugs, has resulted in a renewed focus of the Centers for Disease Control (CDC), health-care
professionals and scientists alike, on multidrug resistant fungal organisms [3, 8]. To add to the problem, *C. auris* reportedly possesses the capacity to adhere to and develop inherently drug-resistant biofilms on abiotic surfaces [10, 28], thus bolstering its ability to persevere in hospital environments, and cause outbreaks. Disseminated infections due to *C. auris* in susceptible patients (e.g. advanced age, presence of central venous catheter, surgery, prolonged hospitalization), are associated with unacceptably high mortality rates between 30–60% [6, 7]. It is unlikely that successful treatment will be achieved with antifungal drugs alone, even with proper antibiotic stewardship. Identification of alternative strategies to prevent and treat infections caused by this fungus is needed. Improved therapy is achievable by either discovering novel molecular targets in its genome, or identifying those that may be homologous to other therapeutically-targeted proteins from phylogenetically related fungi.

We applied computational molecular modeling and bioinformatic strategies to demonstrate that *C. auris* possesses homologs of *C. albicans* major adhesin/invasin protein, Als3p (Fig 1, Table 1 and S1 Dataset). This was an intriguing finding, since Als3p is a hyphae specific protein and *C. auris* does not form filaments. The presence of Als3p-homologs in *C. auris* adds to the list of other genes or gene products with homologies to those associated with virulence and drug resistance in *C. albicans*. For example, homologs of several *C. albicans* efflux genes belonging to the major facilitator superfamily (MFS), MDR and the ATP-binding cassette (ABC) transporter families have been identified, suggesting that efflux is likely a potential resistance mechanism mediating multidrug resistance in *C. auris* [29, 30]. In fact, homologous genes of *C. albicans* virulence proteins such as Serine/Threonine Enzyme-related proteins, mannosyl transferases, a number of secreted aspartyl proteases, as well as kinases involved in virulence and antifungal stress response such as Hog1 protein kinase, 2-component histidine kinase etc., have been identified in the *C. auris* genome [31]. Our discovery of Als3p-like proteins in *C. auris* clearly indicates that despite its extremely high genomic divergence from *C. albicans*, core gene families involved in adhesion/invasion, acquiring drug resistance and other virulence-related genes are conserved in *C. auris*; perhaps also a reason for its persistent nature and its success as a MDR organism.

The Als3p homologs in *C. auris* are targets for vaccine strategies involving the NDV-3A vaccine, which was developed based on the N-terminus region of *C. albicans* Als3, and formulated with alum [18]. Our group has previously reported on the efficacy of NDV-3 (a His-tagged N-terminus of recombinant Als3p formulated with alum) in preventing disseminated candidiasis in mice infected with *C. albicans* [17]. Vaccination with Als3p coupled with complete/incomplete Freund’s adjuvant also protects mice from hematogenously disseminated candidiasis due to other *Candida* species including *C. glabrata*, *C. tropicalis*, and *C. parapsilosis* (S1 Fig). NDV-3A (N-terminus of recombinant Als3p without his-tag and formulated with alum) was also shown to be safe when given to healthy volunteers [18]. In a Phase 1b/2a exploratory clinical trial, NDV-3A was safe and immunogenic in women who suffer from recurrent vulvovaginal candidiasis and protected them from symptoms of infection [19, 20]. Most recently we have described that anti-Als3p antibodies generated from NDV-3A-vaccinated patients (and not alum-treated) had the potential to affect properties of adherence, filamentation and biofilm formation in *C. albicans* [20]. Thus, we questioned if anti-Als3p antibodies could recognize *C. auris*, interfere in functions characteristically associated with Als3p, and prove clinically valuable as vaccine-based strategies against *C. auris*. Using different and complementary approaches (immunofluorescence, flow cytometry, and cell-based ELISA), we validated that NDV-3A vaccination-generated high titer anti-Als3p antibodies that recognized different strains of *C. auris* (Figs 2 and 3 panel A). Importantly, the endpoint IgG antibody titer determined by regular Als3p antigen ELISA and *C. albicans* cell-based ELISA was the same. This finding clearly demonstrates that the two assays are comparable and cell-
based ELISA can correctly estimate the antigen-specific antibody titer in serum. Based on these results, we conclude that although *C. auris* cross-reactive IgG antibodies were 4-fold lower than the Als3p antigen specific IgG antibodies, these titers were still sufficiently high (Fig 3 panel A).

The protective efficacy of NDV-3A vaccine against *C. albicans* infection has been correlated with high antibody titers and Th1/Th17 cell mediated immune responses [17, 21, 23]. An effective immunity against a pathogen is usually comprised of both antibodies and memory T cell immune responses. Therefore, we examined the splenocytes of NDV-3A-vaccinated mice and detected high levels of *C. auris* cross-reactive CD4+ T cells responses in NDV-3A-vaccinated mice, which was similar to Als3p-specific T cells in magnitude. Further, these CD4+ T cells responses (both *C. auris* and Als3p-specific) were marginally shifted towards a Th2-type, probably because alum is known to favor Th2 cell induction [32, 33]. Further, the NDV-3A vaccination also induced high frequency of Als3p-specific Th17 cells that cross-reacted to heat-killed *C. auris*. These results are in line with our previous reports showing that NDV3 and NDV-3A vaccines induce strong Als3p-specific Th17 immune responses [18, 19, 21–23]. Together, these results highlight that NDV-3A vaccine induced robust *C. auris* cross-reactive antibody and T cell immune responses (Fig 3).

The contribution of anti-Als3p antibodies towards abrogation of virulence potential of *C. auris* was illustrated by its ability to block *C. auris* biofilm formation, *in vitro* (Fig 4 panel A). The CAU-09 strain mutant lacking Als3p homolog gene showed reduced binding of anti-Als3p antibodies compared to wild-type CAU-09 strain (Fig 2 panel B). This finding provides evidence that anti-Als3p antibodies obtained from mice vaccinated with NDV-3A, which prevent biofilm formation, indeed bind to the Als3p homologs on *C. auris* surface. Additionally, opsonization of *C. auris* with anti-Als3p antibody containing sera from NDV-3A-vaccinated mice resulted in efficient immune recognition of the pathogen by mice macrophages, leading to increased opsonophagocytic killing (Fig 4 panel B), a feature that was noticed with the activity of anti-Als3p antibodies and *C. albicans* [20, 34]. This finding led to the evaluation of NDV-3A for protection against *C. auris* infections, *in vivo*. Indeed, the results from *in vitro* assays translated successfully *in vivo*, where vaccination by NDV-3A protected some neutropenic mice from disseminated candidiasis caused by *C. auris* (Figs 5 and 6). These preclinical findings suggest that the NDV-3A vaccine may be useful against *C. auris* even in the setting of immunosuppression. Furthermore, although the use of cyclophosphamide in mice results in leukopenia, there is evidence that its effect on tissue macrophages is less severe. In fact, deficiency in alveolar macrophages due to continuous administration of cyclophosphamide was more of a chronic and gradual reduction that never reaches complete deficiency [35]. This finding is in agreement with a study showing that cyclophosphamide has a lesser effect on mouse tissue macrophages, which are usually derived from an embryonic, and not from hematopoietic, origin [36]. Indeed, our mouse macrophage-depletion studies showed that immunosuppressed (cyclophosphamide/cortisone acetate) mice treated with liposome had a considerable number of macrophages in splenocytes (S9 Fig panel B). Thus, the residual tissue macrophages in the cyclophosphamide/cortisone acetate could contribute to the protection seen *in vivo* given the fact that anti-Als3p antibodies enhance OPK activity of macrophages *in vitro* (Fig 4 panel B).

The relative role of humoral and cellular immunity in NDV-3A vaccine-mediated protection against *C. auris* infection was demonstrated by anti-NDV-3A serum adoptive transfer and T cell depletion, respectively (Fig 7). Specifically, our data support a model by which anti-Als3p antibodies are opsonophagocytic and these antibodies and CD4+ cells activate tissue macrophages to enhance fungal clearance. The model is supported by results clearly showing that adoptive transfer experiments with anti-Als3p antibodies can afford protection against *C.
*Candida auris* infection, but are critically dependent on macrophages for efficient protection (Fig 7 panel B). Similarly, depletion of CD4+ compromised the protection afforded by the NDV-3A vaccine (Fig 7 panel C). In this respect, CD4+ T helper cells play a critical role in protecting against fungal infection, and we previously have shown that Th1 and Th17 cells are required for NDV-3A vaccine-mediated protection against *C. albicans* [23].

Aluminum salts together with monophosphoryl lipid A (MPL) are known to prime long-lived memory CD8 T cells [37]. However, NDV-3A vaccine is alum formulated and doesn’t contain any MPL. Thus, it is not expected that NDVA-3A will differentiate CD8+ cytotoxic T-cells. However, the role of CD8+ cell cannot be ruled out especially with the reduced, but not complete reversal of, protection elicited by NDV-3A when CD4+ cells were depleted. In contrast, the complete reversal of protection in macrophage depleted mice argues that the main mechanism of action is reliant on macrophage-mediated opsonophagocytosis clearing of the fungus in infected tissues.

Of great interest and clinical relevance is the additive effect in protecting mice from *C. auris* infection when the vaccine was given with micafungin (Fig 8). This result provides a strong rationale for the continued investigation of the combined use of active NDV-3A vaccination with antifungal drugs in multidrug-resistant lethal *C. auris* infections and gives hope to improved treatment outcomes.

In summary, the unique property of *C. auris* to persist outside human body in the harsh environmental settings of the hospital, its rapid evolution of drug resistance, and its propensity to infect immunosuppressed hospitalized individuals, increasingly threaten global and personal health. The identification of Als3p homologs in *C. auris* opens an avenue for novel immunotherapeutic approaches utilizing either the currently available NDV-3A vaccine, or perhaps an anti-Als3p antibody mediated passive vaccination strategy in the near future. These immunotherapeutic approaches could enhance successful treatment of infections caused by such fungal “superbugs”, thereby reducing morbidity and mortality. Given the safe profile of NDV-3A in humans, future studies will focus on testing this vaccine in patients at high risk of acquiring *C. auris* infection (e.g. colonized patients) and/or those who already suffer from the infection (e.g. therapeutic vaccine combined with antifungal agents).

**Materials and methods**

**Ethical statement**

All procedures involving mice were approved by IACUC of Los Angeles Biomedical Research Institute (protocol 11672), according to the NIH guidelines for animal housing and care. Moribund mice according to detailed and well characterized criteria were euthanized by pentobarbital overdose, followed by cervical dislocation.

**Organisms and culture conditions**

The *C. auris* strains (CAU-01, East Asian clade, ear; CAU-03, African clade, blood; CAU-05, South American clade, blood; CAU-07, South Asian clade, blood; and CAU-09, South Asian clade, bronchoalveolar lavage [BAL]) were obtained from Dr. Shawn Lockhart at Centers for Disease Control and Prevention (CDC, Atlanta). *C. glabrata* 31028, *C. parapsilosis* 22019, and *C. tropicalis* 4243 are all clinical bloodstream isolates obtained from the microbiology laboratory at Harbor–University of California at Los Angeles [UCLA] Medical Center, while *C. krusei* 91–1159 was obtained from the University of Texas Health Science Center at San Antonio. For routine culturing, *C. albicans* (SC5314) and *C. auris* were grown in Yeast Extract Peptone Dextrose (YPD) broth overnight at 30˚C with shaking at 200 rpm. Cells were washed with 1x phosphate buffered saline without Ca**++/Mg**++ (PBS, Gibco by Life Technologies) three times.
prior to counting blastopores with a hemocytometer. For germinating *C. albicans*, 5 \times 10^6 yeast cells/ml in RPMI-1640 media (supplemented with L-Glutamine) were allowed to form germ tubes at 37°C for 75 minutes with shaking at 200 rpm. For staining, cell-based ELISA, and splenocytes stimulation, the *C. auris* strains and Als3p homolog mutants were grown in above germination conditions. For biofilm formation assay, the cells were incubated at 37°C in yeast nitrogen base (YNB) medium for 24 hours.

For heat killing of yeast cells, 5 \times 10^6 cells /mL of PBS were incubated at 65°C for 45 minutes. Yeast cell death was confirmed by plating the heat-subjected cells on fresh plate and incubating the cells at 30°C for several days. For fixation, the yeast cells were incubated with 4% paraformaldehyde solution (in PBS) at 4°C for 1 hour.

**Sequence alignment and analysis**

The *C. albicans* N-terminal Agglutinin-Like Sequence protein (Als3p, Gene Bank ID: AOW31402.1) amino acid sequence was aligned with the *C. auris* using protein BLAST and CLUSTAL-W (NCBI). The predicted Als3p homologs in *C. auris* proteome were further screened for homology among each other using CLUSTAL-W and Als3p homologs having >95% homology with each other considered as same protein. The number of Als3p amino acids showing similarities with *C. auris* proteins was represented as percent positive. The functional domains were identified by using different online bioinformatics tools for GPI-anchor (http://gpi.unibe.ch/), amyloid sequence (http://protein.bio.unipd.it/pasta2/), and Ser/Thr rich sequence (https://www.ebi.ac.uk/Tools/pfa/radar/) prediction. The 3-D protein structure models were built using amino acid sequences and the templates available in the Swiss-model database [38–40] (https://swissmodel.expasy.org/). Briefly, the templates were searched in SWISS-MODEL template library (SMTL) using BLAST and HHHblits. The target sequence was searched with BLAST against the primary amino acid sequence contained in the SMTL. The target-template alignment was performed to build the model by using ProMod3, and coordinates that were conserved between the target and the template were copied from the template to the model. Insertions and deletions were re-modelled using a fragment library, and the side chains were rebuilt. Finally, the geometry of the resulting model is regularized by using a force field. In case loop modelling with ProMod3 fails, an alternative model is built with PROMODE-II. The models showing high accuracy values were finalized for similarity comparisons.

**Preparation of *C. auris* mutants**

To knock-out the Als3p homologs, we generated plasmid constructs containing the nourseothricin resistance gene cassette (pNAT) [41]. For each target Als3p homolog gene, we selected 700–900 bp of the promoter and terminator regions for PCR amplification, as follows: PIS50650.1 gene: upstream fragment (39887–40857) and downstream fragment (45859–46591), PIS50263.1 gene: upstream fragment (923300–924246) and downstream fragment (926539–927376), XP-018167572.2: upstream fragment (849450–8504433) and downstream fragment (852693–853581), (see Table A [I-IV] in S1 Table for primer sequence). We amplified the target gene segments from *C. auris* (CAU-09) genomic DNA and cloned into plasmid pNAT using restriction enzymes (*KpnI/ApaI* for the upstream and *NotI/SacII* for the downstream fragments) or by Gibson method (S1 Fig). Further, to release the gene deletion cassette, the construct was digested with *KpnI/Apal* for the upstream and *NotI/SacII* for the downstream fragments) or by Gibson method (S1 Fig). Further, to release the gene deletion cassette, the construct was digested with *KpnI/SacII* restriction enzymes. The competent *C. auris* (CAU-09) cells were transformed by using gene deletion cassette, CRISPR-Cas9 protein and guide RNA as described previously [42]. Transformants were verified by PCR and RT-PCR using specific primers (Table A in S1 Table) to confirm the deletion of the target gene [43] (S1 and S2 Figs).
Immunization of mice

We used 4–6 week old outbred ICR (CD-1) mice in this study. The NDV-3A vaccine was formulated by mixing 300 μg of *C. albicans* recombinant Als3p with 200 μg alum adjuvant per dose. The recombinant Als3p was produced in *Saccharomyces cerevisiae*, and was a Gift from NovaDigm therapeutics. For immunization, the NDV-3A vaccine or alum was injected into mice subcutaneously (s.c.) at day 0, 21 and 35. Mice were bled 14 days after final immunization and serum was isolated for antibody titer determination. For efficacy studies against *Candida* species other than *C. auris*, mice were immunized by subcutaneous injection of recombinant N-terminus of Als3p-N (20 μg) mixed with complete Freund’s adjuvant (CFA; Sigma-Aldrich, St. Louis, MO) at day 0, followed by a booster dose in incomplete Freund’s adjuvant (IFA; Sigma-Aldrich) at day 21. Control mice were immunized with CFA followed by IFA alone. Fourteen days following the boost, immunized mice were infected via the tail vein with the inocula mentioned in S11 Fig.

Confocal microscopy and flow cytometry

The yeast or germ tube cells (2x10^6 cell) from wild type *C. auris* strains (CAU-01, CAU-03, CAU-05, CAU-07 and CAU-09), CAU-09 Als3p homolog mutants (PIS50650.1-/-, PIS50263.1-/-, XP-018167572.2-/-) or *C. albicans* were fixed with 4% paraformaldehyde at 4˚C for 1 hour. After blocking the cells with 3% bovine albumin serum solution (in 1x PBS), these cells were added to 96-well plate and centrifuged to pellet the cells. The pellet was resuspended in 100 μl of anti-NDV-3A or alum serum diluted at 1:500 in 1x PBS and incubated for 1 hour at room temperature. The cells were washed three times with 1x PBS prior to adding 100 μl of Alexa Fluor 488 labelled anti-mouse IgG detection antibodies (1:100 dilution in PBS). After 1 hour of incubation at room temperature, the cells were resuspended in 300 μl of PBS and analyzed using confocal microscopy or flow cytometry.

For confocal microscopy, 20 μl of stained cells were added to the glass slides and covered with cover slips. The images were taken at 40x resolution using Laser Scanning Confocal microscopy (Leica Confocal Microsystem). For flow cytometry, the stained cell suspension was transferred to flow tubes and 20,000 events were acquired using LSR II flow cytometer (BD Biosciences). Flow cytometry data were analyzed using FlowJo software (Version 10).

Enzyme-linked immunosorbent assay

Ninety six-well plates were coated with 5 μg/ml of Als3p in bicarbonate/carbonate coating buffer (pH 9.6) overnight at 4˚C. The next day, the plates were washed three times with 1x wash buffer (PBS containing 0.05% tween-20) and blocked with 3% BSA solution for 2 hours at room temperature. After washing three times, diluted serum samples were added to the plates in duplicates and incubated for two hours. After incubation, the plates were washed three times and 1:1000 diluted anti-mouse IgG antibodies (Jackson, Cat#115-035-164) labelled with peroxidase were added and incubated for 1 hour at room temperature. Finally, the plates were washed five times with washing buffer, TMB (3,3',5,5'-Tetramethylbenzidine) substrate (Invitrogen, Cat#00-4201-56) was added. Color development was allowed for 5–10 minutes and absorbance was measured at 450 nm after the reaction was stopped with 1 N sulfuric acid (Sigma, Cat#339741).

To determine the *C. auris* cross-reactive antibodies in NDV-3A vaccinated mice, we used a cell-based Enzyme-Linked Immunosorbent Assay (ELISA). As above, germinated *C. albicans* (SC5314) or *C. auris* (CAU-09) were counted and fixed in 4% cold paraformaldehyde (Sigma-Aldrich, Cat#158127) for 30 min. The fixed cells were washed with 1x PBS. The U-bottom 96-well plates were blocked overnight with 1x blocking solution (3% BSA in 1x PBS, Thermo...
Fischer), and next day 2-fold serial dilutions of 100 μl serum samples (diluted in 1x blocking solution) per well were added in duplicates. Further, a total of 10^7 fixed cells/well/100 μl of C. auris and germinated C. albicans were added to these serum sample-containing wells. The plates were incubated for 2 hours at room temperature. After incubation, the cells in the 96-well plate were washed three times with 1x PBS and secondary HRP-conjugated anti-mouse IgG antibodies were added as per the manufacturer instructions. After 1 hour incubation at room temperature, the cells in 96-well plate were washed with 1x PBS. Further, 100 μl/well TMB substrate was added to each well and incubated until the color developed (normally 5–20 min). The color reaction was stopped by adding 50 μl 1 N H₂SO₄ per well and plates were centrifuged at maximum speed to pellet the cells. One hundred microliter colored substrate supernatant from each well were transferred to fresh flat bottom 96-well plates and OD was measured at 450 nm. The endpoint titers in NDV-3A vaccinated mice were determined by plotting mean OD450 vs serial serum dilution, and noting the highest dilution with significantly higher OD450 compared to alum vaccinated mice group.

**Intracellular cytokine staining and flow cytometry**

The splenocytes were harvested from the NDV-3A or alum immunized mice by homogenizing individual spleens in 100 μm cell strainer. The RBCs were lysed by 1x RBC lysis buffer (Santa Cruz Biotech, Dallas, Cat# SC-296258), and filtered through the 100 μm sterile filters. The cells were resuspended in 1x RPMI supplemented with 10% FBS, counted and plated at 1x10^6 splenocytes/100 μl/well in a U-bottom 96-well plate. The splenocytes were kept unstimulated or stimulated with 100 μl/well of 10 μg/ml recombinant Als3p, or 3x10^6 cells/well heat-killed C. auris for 5 days at 37˚C. On day 5, 100 μl of the culture supernatant was replaced with fresh RPMI media containing 50 ng/ml PMA (phorbol 12-myristate 13-acetate, Sigma, Cat# P8139), and 500 ng/ml ionomycin (Sigma, Cat#I3909). After 1 hour, protein transport inhibitor cocktail (eBioscience, Cat# 00-4980-03) was added to each well at final 1x concentration, and plates were incubated for another 3 hours. Next, the cells were stained with extracellular CD3 APC-Cy7 (BD Biosciences, Cat#557596) and CD4 PerCP Cy5.5 (BD Biosciences, Cat#550954) antibodies (0.25 μg/sample) for 30 minutes at room temperature. Subsequently, the cells were washed, fixed and permeabilized with Cytofix/Cytoperm solution (Invitrogen, Cat#00-5123-43) and then stained intracellularly with IFN-γ APC (BD Pharmingen, Cat#554413), IL-4 FITC (eBioscience, Cat#11-7042-82) and IL-17 PE (R&D Systems, Cat#IC7211P) antibodies (0.25 μg/sample) for 30 minutes at room temperature. The stained cells were acquired in BD LSR-II flow cytometer (BD Biosciences), and at least 30,000 CD3+ T cells were recorded. The data was analyzed using FlowJo (Version 10) software.

**Biofilm formation assay**

Biofilms were developed in 96-well polystyrene microtiter plates as previously described, with slight modifications [44]. Briefly, 95 μL of C. auris cells (2 × 10^5 cells/ml in YNB medium) was added to the wells containing 5 μL of test or control mouse serum (5% serum vol/vol), and incubated at 37˚C. Control wells had no serum. After 24 hours, wells were washed twice with PBS, and the extent of biofilm formation was quantified by XTT assay (490 nm) [44].

**Opsonophagocytic killing (OPK) assay**

The opsonophagocytic killing assay was based on a modification of a previously used method [45]. C. auris yeast cells (CAU-09) were added into 96-well microtiter plates. Murine RAW 264.7 macrophage cells (American Type Culture Collection [ATCC# TIB-71], Rockville, MD) were cultured at 37˚C in 5% CO₂ in RPMI-1640 (Irvine Scientific, Santa Ana, CA) with 10%
fetal bovine serum (FBS), 1% penicillin, streptomycin, and glutamine (Gemini BioProducts), and 50 mM β-mercaptoethanol (Sigma-Aldrich, St. Louis, MO). RAW 274.7 cells were activated by exposure to 1 ng/ml LPS (Sigma-Aldrich) for 24 hours. Activated RAW 264.7 macrophages were harvested after scraping with BD Falcon cell scrapers (Fischer Scientific) and added to the microtiter wells at a 1:1 ratio of macrophages to \( C. auris \). After 2 hours of incubation with gentle shaking, aliquots from the wells were quantitatively plated in YPD agar plates. The percent killing of \( C. auris \) was calculated using the following formula: 1 - [CFUs from tubes with (mouse serum + \( C. auris \) + macrophages) /average CFU in tubes with (\( C. auris \) + macrophage)].

Mice infection and treatment

CD-1 mice vaccinated with NDV-3A vaccine were infected with \( C. auris \) CAU-09 to evaluate the efficacy of the vaccine. Briefly, twelve days after the final boost, mice were made neutropenic by 200 mg/kg cyclophosphamide delivered intraperitoneally (i.p.) and 250 mg/kg cortisone acetate (s.c.) administered on days -2 and +3, relative to infection. To prevent bacterial superinfection in the immunosuppressed mice, we added enrofloxacin (at 50 μg/ml) to the drinking water. Mice were infected through tail vein injection with \( 5 \times 10^7 \) CFU/mouse. For combination with antifungals, alum- or NDV3-A-vaccinated and infected mice were treated with a minimal protective dose of 0.5 mg/kg/day of the clinically used micafungin by i.p. administration. Treatment started after 24 hours of infection and continued until day +7. Mice were monitored for their survival for 21 days after the infection.

For fungal burden determination, mice were vaccinated, made neutropenic and then infected as above and then euthanized on day 4 post infection to collect kidneys and brain. The organs from each mouse were weighed, homogenized and quantitatively cultured by 10-fold serial dilutions on YPD plates. Plates were incubated on 37°C for 48 hours prior to enumerating colony forming units (CFUs)/gram of tissue. Finally, histopathological examination of kidneys or brain from mice sacrificed on Day 4 post infection, were fixed in 10% zinc-buffered formalin, embedded in paraffin, sectioned, and stained with Pacific Acid Schiff (PAS) stain.

Serum adoptive transfer and immune cell depletion experiment

Balb/c mice were vaccinated with NDV-3A or alum on day 0, and boosted on days 21 and 35. Vaccinated mice were made neutropenic using 200 mg/kg body weight of cyclophosphamide administered subcutaneously and 250 mg/kg body weight of cortisone acetate given by intraperitoneal injection on day 47. Mice were infected intravenously with \( C. auris \) on day 49 as above. To prevent bacterial superinfection in the immunosuppressed mice, 5 mg ceftazidime /mouse was administered daily through subcutaneous route for up to 14 days post infection, and after that 50 μg/ml enrofloxacin was added to drinking water. All NDV-3A and alum vaccinated mice were grouped as depletion and control depletion arms. For CD4+ T cell depletion experiment, 200 μg/mouse dose of rat anti-mouse CD4 IgG2b (clone GK1.5, BioXcell, Cat BE0003-1) or rat IgG2b isotype antibodies (Clone: LTF-2, BioXcell, Cat #BE0090) were administered intraperitoneally on day 11 and 14 post-second boost (i.e. days -3 and 0, relative to infection) [27, 46]. For macrophage depletion, 1 mg clodronate-liposome or PBS-liposome particles (Liposoma B.V., Cat # CP-005-005) were injected intraperitoneally on day 11 and 14 post second boost, and subsequently twice a week (i.e. days -3 and 0, 4, 8, 12 and 16 relative to infection) [47, 48]. Mice were infected intravenously with \( 5 \times 10^7 \) \( C. auris \) (CAU-09) and monitored for their survival for three weeks. Three additional mice in each depletion and control depletion arm were taken to verify the CD4 and macrophage depletion. Briefly, 4 days after administering the second dose of
depletion drug, mice were euthanized, and spleen and lymph nodes were harvested and homogenized to make single cell suspension. For CD4+ cell depletion verification, splenocytes and lymph node cells were stained with anti-CD3 APC (BD Pharmigen, Cat #BDB565643) and anti-CD4 Alexa Fluor 700 antibodies (Biolegend, Cat #100536). Macrophage depletion was verified by staining anti-CD11b PerCP Cy5.5 (eBioscience, Cat #5015759) and anti-F4/80 APC (eBioscience, Cat #501128925) antibodies. Stained cells were acquired in BD LSR II flow cytometer, and data were analyzed in FlowJo software.

Statistical analysis

Differences in survival studies were analyzed by the non-parametric Log Rank test for overall survival and with Mantel-Cox comparisons for median survival times. All other comparisons were conducted with the non-parametric Mann Whitney test. P values <0.05 was considered significant.

Supporting information

S1 Dataset. Protein sequence alignment. (A) Comparison between C. albicans Als3p (GenBank: AOW31402.1) and its homolog on C. auris (GenBank: PIS50650.1) using CLUSTAL-W. (B) Comparison between C. albicans Als3p (GenBank: AOW31402.1) and its homolog on C. auris (GenBank: PIS50263.1) using CLUSTAL-W. (C) Comparison between C. albicans Als3p (GenBank: AOW31402.1) and its homolog on C. auris (GenBank: XP_018167572.2) using CLUSTAL-W.

S1 Fig. Schematic of the gene deletion construct using nourseothricin (NAT1) resistance cassette. (A) Preparation of the gene deletion construct using restriction enzymes. A-1. DNA sequences of the promoter (upstream fragment) and terminator (downstream fragment) regions of the target gene were PCR-amplified. A-2. The upstream fragment was digested by KpnI and Apal restriction enzymes and cloned into the NAT1 cassette. A-3. The downstream fragment was digested by NotI and SacII restriction enzymes and cloned into the NAT1 cassette. A-4. The competent C. auris (CAU-09) cells were transformed by gene deletion cassette, guide RNA and Cas9 protein by using electroporation. (B) The upstream fragment was cloned into NAT cassette by using PCR and the downstream fragment was digested by NotI and SacII restriction enzymes and cloned in NAT1 cassette.

S2 Fig. PCR and RT-PCR analysis of transformants to detect the null mutants of each gene. (A) PCR amplification of PIS50650.1, PIS50263 and XP-018167572.2 genes was performed using a set of primers for each gene (Table A in S1 Table). For PIS50650.1, colony # 3 and 5, for PIS50263, colony # 2 and for XP-018167572.2, colony # 4 did not show a PCR band. (B) RT-PCR was done for PIS50650.1 # 3, and PIS50263 # 2 and XP-018167572.2 # 4 colonies using specific primers (Table A in S1 Table) to study the gene expression of each gene at mRNA level compared to CAU-09 as wild type. M = DNA marker (bp).

S3 Fig. Intracellular cytokine staining gating strategy and Th1 (CD3+ CD4+ IFN-γ +), Th2 (CD3+ CD4+ IL4+) and Th17 (CD3+ CD4+ IL17+ phenotype determination.

S4 Fig. C. auris cross-reactive Th1 (CD4+ IFN-γ +) cells in NDV-3A vaccinated mice.
S5 Fig. *C. auris* cross-reactive Th2 (CD4+ IL4+) cells in NDV-3A vaccinated mice. (TIF)

S6 Fig. *C. auris* cross-reactive Th17 (CD4+ IL17+) cells in NDV-3A vaccinated mice. (TIF)

S7 Fig. Sera from NDV-3A or alum vaccinated mice have no effect on viability of *C. auris*. (TIF)

S8 Fig. Anti-NDV-3A pooled immune sera preparation and anti-Als3p IgG antibody titer determination. (TIF)

S9 Fig. Macrophage depletion verification in mice treated with clodronate-liposome. (TIF)

S10 Fig. CD4+ T helper cell depletion verification in mice treated with anti-CD4 (GK1.5) antibodies. (TIF)

S11 Fig. Recombinant N-terminus Als3p-based vaccine showed reduced fungal burden in mice infected with several non-albicans species of Candida: *C. glabrata*, *C. krusei*, *C. parasilosis*, or *C. tropicalis*. Immunocompetent BALB/c mice (n = 5 per group) were vaccinated with 20 μg N-terminus Als3p/mouse mixed with complete Freund’s adjuvant (CFA) on Day 0 then boosted with similar dose mixed with incomplete Freund’s adjuvant (IFA) on Day 21 and infected via the tail vein on Day 34. Infectious inocula are shown in parentheses below the species names. Kidney fungal burden was determined on day 5 post-infection. *p < 0.05 versus CFA control by using nonparametric Steel test for multiple comparisons*. (TIF)

S1 Table. (A)Primer and guide RNA sequence used in Als3p homolog mutant preparation. (B) MIC of antifungal drugs against different *C. auris* strains. All MIC are in μg/ml and determined by CLSI method. For azoles and echinocandins, MIC is defined as 50% inhibition, and for amphotericin B as 100% inhibition of different *C. auris* strains. (DOCX)

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