Chickens with humanized immunoglobulin genes generate antibodies with high affinity and broad epitope coverage to conserved targets

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ABSTRACT
Transgenic animal platforms for the discovery of human monoclonal antibodies have been developed in mice, rats, rabbits and cows. The immune response to human proteins is limited in these animals by their tolerance to mammalian-conserved epitopes. To expand the range of epitopes that are accessible, we have chosen an animal host that is less phylogenetically related to humans. Specifically, we generated transgenic chickens expressing antibodies from immunoglobulin heavy and light chain loci containing human variable regions and chicken constant regions. From these birds, paired human light and heavy chain variable regions are recovered and cloned as fully human recombinant antibodies. The human antibody-expressing chickens exhibit normal B cell development and raise immune responses to conserved human proteins that are not immunogenic in mice. Fully human monoclonal antibodies can be recovered with sub-nanomolar affinities. Binning data of antibodies to a human protein show epitope coverage similar to wild type chickens, which we previously showed is broader than that produced from rodent immunizations.

Introduction
Monoclonal antibody (mAb) therapy is a crucial segment of the pharmaceutical arsenal, with 57 FDA-approved mAbs, an estimated nine mAbs predicted to be granted their first marketing approvals in 2017, and hundreds more in development. Therapeutic mAbs should be as similar to native human antibodies as possible to minimize their immunogenicity in patients. Historically, human therapeutic mAbs in the clinical pipeline have been produced from three main technologies: 1) in vitro complementarity-determining region (CDR) grafting of murine antibodies onto human frameworks, 2) in vitro systems such as phage display libraries, and 3) in vivo immune systems of “humanized” mice genetically engineered to express a human immunoglobulin repertoire. To date, the majority of approved human mAbs have been derived from the mouse (wild type (WT) or transgenic) rather than in vitro systems. Antibodies produced in the intact immune system of an animal have gone through rigorous selection for specific binding to the target, counter-selection to a vast array of endogenous off-target proteins, and high-level expression in plasma cells. The in vitro-derived antibodies do not have the advantage of immune system selection, and therefore require further validation and sequence optimization after initial discovery.

The strategies for expression of human antibodies in transgenic animal platforms have evolved over time, and the range of species that have been genetically modified has expanded to include mice, rats, rabbits, and cows. As all these transgenic animals are mammals, the accessible epitope space on human protein targets is limited. Immune responses are primarily focused on regions that are evolutionarily divergent between the species, and it is difficult to raise antibodies against epitopes that are shared amongst mammals, i.e., “pan-mammalian” epitopes, in a mammalian host. Nonetheless, species cross-reactive antibodies are highly desirable for drug development because such antibodies facilitate the use of animal models of disease. When a lead therapeutic antibody is not species cross-reactive, it often necessitates a parallel campaign for a “surrogate” antibody.

As an alternative to mammalian species, birds (and in particular, chickens) present an attractive choice because they are phylogenetically distant from humans (having diverged about 300 million years ago). Transgenic animal models of disease have evolved over time, and the range of species that have been genetically modified has expanded to include mice, rats, rabbits, and cows. As all these transgenic animals are mammals, the accessible epitope space on human protein targets is limited. Immune responses are primarily focused on regions that are evolutionarily divergent between the species, and it is difficult to raise antibodies against epitopes that are shared amongst mammals, i.e., “pan-mammalian” epitopes, in a mammalian host. Nonetheless, species cross-reactive antibodies are highly desirable for drug development because such antibodies facilitate the use of animal models of disease. When a lead therapeutic antibody is not species cross-reactive, it often necessitates a parallel campaign for a “surrogate” antibody.

As an alternative to mammalian species, birds (and in particular, chickens) present an attractive choice because they are phylogenetically distant from humans (having diverged about 300 million years ago).
years ago, compared to 65 million years for rodents), produce antibodies of high affinity and specificity, and can recognize unique epitopes not accessible in mice. Expanded epitope coverage is an advantage in drug development because it increases the chances of accessing the functionally significant regions of the target, which would likely be conserved. Antibodies with unique characteristics may be identified that provide novel mechanisms of action. Cross-reactive mAbs that recognize the human and mouse orthologs of a target could seamlessly advance from pre-clinical to clinical-stage studies. These benefits are achieved while retaining the general advantage of an in vivo selection process that removes non-specific and poorly expressing clones.

Chicken display broad epitope coverage and can generate served proteins that are not immunogenic in mice. The OmniChicken retains the expanded epitope coverage observed in WT chickens, and frameworks normally found in chickens. The OmniChicken the chicken immune system and to take advantage of the restricted species, such as mice and cynomolgus monkeys, that are relevant to mechanism-of-action and toxicology pre-clinical studies. The human transgenes have been designed to work in the context of

The complex genetic modifications necessary to make the OmniChicken were produced in cultured germline cells, which were then used to obtain fully transgenic chickens. The immunoglobulin loci of the chicken were modified in a two-step process: 1) targeting of an attP site into the light and heavy chain loci by homologous recombination, then 2) insertion of human sequences using phiC31 integrase (Supplementary Fig. 1). The attP insertion step simultaneously deleted endogenous Ig sequences, producing gene knockouts. Phenotypic analysis of the knockouts confirmed that the correct loci were targeted and that Ig expression was eliminated. In both light and heavy chain knockouts, the endogenous upstream pseudogenes remained intact. In the second step, single functional human VH and VK genes were inserted site-specifically into the attP sites targeted to the Ig loci and were designed to splice to the endogenous chicken constant regions, to ensure proper formation of the B cell receptor complex and engagement of chicken Fc receptors (Supplementary Fig. 1).

Here we present the OmniChicken, a transgenic chicken carrying humanized immunoglobulin genes that can be used to discover novel, high affinity antibodies, including antibodies against conserved proteins that are not immunogenic in mice. The OmniChicken displays broad epitope coverage and can generate antibodies that are cross-reactive with homologs in mammalian species, such as mice and cynomolgus monkeys, that are relevant to mechanism-of-action and toxicology pre-clinical studies. The human transgenes have been designed to work in the context of the chicken immune system and to take advantage of the restricted frameworks normally found in chickens. The OmniChicken retains the expanded epitope coverage observed in WT chickens, but in conjunction with human-sequence antibodies.

**Results**

Peripheral blood mononuclear cells (PBMCs) from WT, homozygous heavy-chain knockout (IgH KO), and OmniChickens were labeled with antibodies to chicken IgM and IgL constant regions, the B cell marker Bu-1, and T cell receptors, and analyzed by flow cytometry. In the IgH KO birds, all B lineage cells are missing, and in the OmniChickens, the B cell population was reconstituted to normal levels (Fig. 1A). The expression level of surface IgM on circulating cells was normal, and only expressed on B cells (Fig. 1A). The percentages of B and T cells in the periphery were normal (Fig. 1B).
Plasma levels of IgM, IgY and IgA

Chickens contain three immunoglobulin isotypes: IgM, IgY and IgA. IgY is functionally analogous to mammalian IgG, although it contains four CH domains and no hinge. As in mammals, IgM is the first expressed isotype in chickens and class switching leads to expression of IgY or IgA. IgA is mainly found in bile and other secretions, but is also found in plasma. Plasma levels of all three isotypes were analyzed by ELISA (Fig. 2). Whereas plasma levels of IgA were normal, levels of plasma IgM and IgY were found to be reduced (3-fold and 15-fold, respectively). The lower relative levels of IgM and IgY were consistent from 4 weeks to 28 weeks (adulthood). In birds expressing a human V-lambda light chain (instead of the

![Figure 1](image1.png)

**Figure 1.** Flow cytometry analysis shows normal B cell development in the OmniChicken and normal levels of surface IgM. A, Wild-type, IgH knockout (IgH KO), and OmniChicken PBMCs were labeled with antibodies against the B cell marker Bu-1 and either IgL or IgM (left), or with antibodies against IgL and IgM (right). B, Flow cytometry analysis of lymphocytes shows normal levels of B and T cells. Wild-type (N = 3 birds) and OmniChicken (N = 12 birds) PBMCs were labeled with antibodies for Bu-1, IgL, IgM and T cell markers TCR1 and TCR2/3.

**Plasma levels of IgM, IgY and IgA**

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![Figure 2](image2.png)

**Figure 2.** Plasma levels of IgM and IgY are reduced in OmniChickens but IgA levels are normal. Immunoglobulins were measured by ELISA in wild-type (N = 7) and OmniChickens (N = 9). Ig concentrations were determined by comparison to standards. Mean values (in μg/mL) and standard deviations are shown below the plots.
V-kappa in the OmniChicken), IgY levels were normal (Supplementary Fig. 3), suggesting that the V-kappa light chain is somehow limiting antibody plasma levels in the OmniChicken.

**Immunization and cloning of human mAbs**

OmniChickens were immunized with a variety of proteins to investigate whether the transgenic birds retained the advantages of the chicken host system, while producing fully human antibodies to a wide range of epitopes on both conserved and non-conserved targets. As a first test of immune competence, OmniChickens were immunized with a cocktail of antigens (bovine thrombin, human transferrin, and murine IL-13). Antigen-specific titer was observed in plasma from birds for all of the antigens (Fig. 3A), which increased after subsequent boosts. After reaching an appropriate titer, spleens were obtained and lymphocytes prepared for mAb recovery.

A robust hybridoma fusion partner is not available in chickens, so we have developed a single B cell screening method called the GEM assay (for Gel Encapsulated Microenvironment) that can be used to identify antigen-specific antibody-secreting cells from any source, including OmniChickens.10,13 Spleen cells are encapsulated in agarose droplets along with polystyrene beads that have been coated with the target antigen. After incubation to allow secretion of IgY antibody and binding to the antigen on the beads, the IgY is detected with a fluorescent secondary antibody. Bead-positive GEMS are viewed and selected at the microscope, and single cell RT-PCR is used to amplify and clone the variable regions. The V region amplicons from each positive GEM are joined together by a linker sequence to generate a single-chain variable fragment (scFv), and then cloned into a vector containing a human Fc. This expression cassette thus produces a fully human recombinant antibody in the bivalent scFv-Fc format.

GEM screens were performed individually for each of the cocktail antigens and target-specific mAbs were recovered from individual birds for each antigen in the cocktail. These data confirm the ability of the OmniChicken to produce antigen-specific mAbs (Fig. 3A).

Array-based surface plasmon resonance imaging (Array SPRi) was used to perform epitope binning experiments on 8 of the thrombin-specific antibodies. These 8 clones clustered into 3 distinct non-overlapping bins (Fig. 3B). Sequence analysis (Fig. 3C) showed that the 5 mAbs in bin A are highly related, whereas the 2 mAbs in bin C are more distantly related, indicating that diverse sequences can converge on the same epitope.

To test more rigorously the ability of OmniChickens to recognize highly-conserved targets, OmniChickens were immunized with human brain-derived neurotrophic factor (BDNF), a protein that is 97% conserved between humans and mice, and 91.5% conserved between humans and chickens. The response to BDNF was variable. Of 9 birds immunized with BDNF protein, 2 had no response, 3 had low titers of under 1:500, but 4 birds had good responses, with titers of 1:4,050 to 1:24,300 (Fig. 4A). Spleens were harvested from the immunized birds, and mAbs recovered using the GEM assay. Epitope binning using Array SPRi showed that all six clones mutually blocked one another, and therefore were assigned to the same epitope bin. This is not surprising given the relatively small surface-accessible area of this homodimer (total 28 kDa) that is not buried at the dimer interface.8,13 It is striking that the six anti-BDNF mAbs display a high level of sequence diversity, particularly in the heavy chain, despite all converging on the same epitope bin (Fig. 4B). This result indicates that it is not possible to state whether antibodies with highly divergent sequences will necessarily bind their target differently; unrelated clones can converge on the same epitope, especially if the available epitope space is restricted.

To further characterize their antibody repertoire, OmniChickens were immunized with human progranulin (PGRN), a multi-domain protein we have used previously to interrogate epitope space covered by WT chickens as compared to WT mice.8,14 Birds were either immunized with human PGRN protein or DNA encoding human PGRN. OmniChickens produced robust titers to PGRN protein (up to 1:36,450), similar to those obtained in WT birds.8,14 A panel of mAbs was produced from both protein and DNA-immunized OmniChickens by screening in GEMs using human PGRN-coated beads.

Array SPRi was used to perform epitope binning on a panel of >100 PGRN mAbs from OmniChickens (Fig. 5). All 7 of the

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**Figure 3.** Individual OmniChickens raise specific mAbs to multiple targets. Birds were immunized with a cocktail of mouse IL-13, human transferrin and bovine thrombin, human transferrin, and mouse IL-13. A, Plasma titers to each immunogen from a single bird are shown over multiple draws (top set of graphs). Panels of mAbs to each antigen were identified from that bird, and antigen-specific binding of a single representative mAb for each target is shown (bottom set of graphs). B, Epitope binning of eight thrombin antibodies shows three distinct bins. C, Phylogenetic tree of the 8 thrombin antibodies, with the epitope bin indicated at right.
granulin domains were recognized by members of the panel, and some members of the panel recognized human/mouse cross-reactive epitopes, thereby mirroring the epitope diversity we reported previously for their WT counterparts. Kinetic analysis showed a range of binding affinities from 0.11 to ~200 nM (mean value, 25.6 nM), including some in the sub-nanomolar category, although the highest affinities were found in mAbs from WT birds (the highest being 10 pM). Sequence analysis showed that diverse sequences can converge on the same epitope bin.

**Sequence analysis**

Sequences of the VK and VH regions were determined for all of the antigen-specific antibodies described above, and the diversity across the panels of antibodies at each position is shown in Supplementary Figures 4 and 5. All of the clones showed accumulation of mutations in both heavy and light chain V regions relative to the original (germline) transgene sequence. Somatic mutations were concentrated in the CDRs, with CDR-H3 showing the most diversity. Gene conversion using sequences...
donated by the synthetic human pseudogenes was observed in both the VK and VH variable regions, as well as point mutations that cannot be ascribed to gene conversion. All of the human pseudogenes were used in gene conversion; no sequences derived from the chicken pseudogene pool were observed for either light or heavy chain. Sequences of mAbs from different birds were usually quite divergent, although we observed several examples of related sequences from different PGRN-immunized birds that were a result of clear gene conversion leading to similar CDRs being incorporated (Fig. 5D shows one such example). These related antibodies also converged on the same epitope bins. In addition, some residues were commonly found to be introduced by gene conversion into a majority of the antibody sequences. The Arg-Leu-Phe (RLF) motif present in heavy chain framework 3 (IMGT positions 90–92) in the functional V of the SynVH-C construct was gene converted in virtually every antibody sequence to Gln-Met-Asn (QMN) (Supplementary Fig. 4), which can be found in several of the pseudogenes in the SynVH-C array. A Gly residue in CDR-L2 (IMGT position 56) was found to be changed to Asp in about half of the antibody sequences from multiple birds immunized with PGRN. For the antibodies derived from the SynVH-A7 array (Supplementary Fig. 5), in which heavy chain pseudogenes contain only Ser, Trp and Tyr residues in the CDRs, heavy chain gene conversion was more limited than in SynVH-C and consisted of shorter stretches of sequence encoding only one or two amino acids. The longest gene conversion tract was found in CDR-H3 where the residue pair Ser→Tyr was found in a subset of mAbs and was likely derived from pseudogene SynVH17. The majority of the heavy chain diversity in SynVH-A7 mAbs consisted of sequences not found in the pseudogenes.

Some variation in CDR-H3 length was observed, with a range of 9 to 16 amino acids. A substantial proportion (40%) of the sequences contained 13 amino acids in CDR-H3; the germline pre-rearranged CDR-H3 contained 11 amino acids. No cysteine residues were found in any of the CDR-H3 sequences, in contrast to WT chickens, which often contain cysteine in CDR-H3.92 (27/32 clones in a panel of anti-PGRN chicken mAbs37). The only non-canonical cysteine that was found in the human sequences was in one clone (out of ~150) that had an unpaired cysteine in VH framework 2.

Discussion

Human antibody therapeutics have become an integral part of the pharmaceutical space, and for many indications are a preferred drug format due to their high target-specificity, effector functionality and long serum half-life. During the process of discovery and development, a variety of attributes need to be examined in candidate antibodies, including biophysical properties, immunogenicity, toxicity, efficacy, potency, and pharmacokinetics. Many areas of evaluation require the use of animal models of disease, most commonly in rodents or non-human primates. Therefore, it is advantageous to source antibodies from a phylogenetically distant species that will recognize the human target and its orthologs as foreign and readily elicit antibodies to pan-mammalian epitopes. Chickens have long been used to raise such antibodies against conserved mammalian targets.28–33 Here, we describe an engineered transgenic chicken that retains native antigen recognition, but delivers human sequence antibodies that are suitable for therapeutic use. Replacing the chicken V gene repertoire with a human one allows the transgenic bird to generate human antibodies that retain the benefits of the chicken host, namely broad epitope coverage of the target beyond that available from mammals. The expanded epitope coverage includes a prevalence of species cross-reactive antibodies, which in many cases obviates the need for surrogate antibodies. Targeting novel epitopes can also provide opportunities for therapeutics with different mechanisms of action than previously available and expand the possible routes to drug development. We have shown that the OmniChicken, like WT birds, can recognize highly conserved mammalian proteins such as BDNF, and can be used to provide panels of unique antibodies of high affinity to multiple epitopes on human targets.

The B cells in the OmniChicken are capable of producing a high level of sequence diversity in the human VK and VH genes using both gene conversion and non-templated mutation. In the heavy chain, two different strategies were pursued to produce diversity. In SynVH-C, the pseudogenes contained diverse CDRs and limited diversity in the frameworks, all sourced from naturally occurring human sequences. The power of in vivo selection was evident in that certain sequences, such as the QMN in framework 3, could be obtained from the pseudogene pool and positively selected. In SynVH-A7, the pseudogenes contained a restricted set of residues in the CDRs (only Y, W and S), whereas the CDRs found in the antibody sequences were much more diverse, indicating that additional amino acids were incorporated by somatic hypermutation. Both constructs yielded sequences that could be ascribed to gene conversion, with somatic hypermutation and selection adding another layer of diversity in the CDRs and frameworks. In both the light and heavy chain loci, the chicken pseudogenes are still present upstream of the human transgenes. We saw no evidence of chicken sequence being introduced into the human V regions (references 34 and 35 and Supplementary Figs. 4 and 5). Gene conversion is a homology-based process, and the human V pseudogenes have much higher homology to the human functional V, and they are in closer proximity, which could introduce further bias in their favor.36 Although the SynVK and SynVH transgenes contain fewer pseudogenes than the WT loci (16 vs. 25 for the light chain and 20 vs. 100 for the heavy chain), not all of the chicken pseudogenes may participate in gene conversion.36 In contrast, all of the human pseudogenes were used in our panels and in previous work in the DT40 model.35 The overall level of sequence diversity in the cohorts of anti-PGRN mAbs derived from OmniChicken and WT birds seemed similar when comparing the sequence dendrograms.8 Most importantly, the repertoire diversity generated by gene conversion or point mutation in OmniChicken and WT birds recognized equivalent epitope space on the PGRN protein.

The OmniChicken expresses chimeric (human variable/chicken constant) immunoglobulins, to ensure the proper engagement of endogenous constant region receptors and to form the B cell receptor complex with Igα/β.37 Once the human V regions are identified by cloning, they can be combined with any Fc region or into any antibody format using standard
cloning techniques. Typically, our mAbs are assembled in the scFv-Fc format for initial evaluation. Subsequently, the best candidates are reformatted into full length IgG molecules with equivalent specificity, sensitivity and affinity constants. In addition, they are expressed at normal levels in mammalian cell culture.

Levels of plasma Ig were normal or near normal for IgM and IgA isotypes, but lower for IgY. Reduced Ig levels have also been observed in many of the humanized rodent platforms.\(^5\) Circulating IgY is produced by terminally-differentiated plasma cells, which contain a greatly expanded secretory apparatus for high-level secretion of antibody. It is possible that either the number of plasma cells is reduced, or that IgY expression per cell is lower than normal. In chickens, markers to specifically assess the number of plasma cells are lacking, but the numbers of splenocytes from immunized OmniChickens were comparable to those from immunized WT birds (2.2e9 splenocytes averaged over 25 OmniChickens compared to 2.5e9 splenocytes averaged over 28 WT birds). Reduced expression of the human kappa chimeric light chain in IgY-switched plasma cells seems to be the main cause of the IgY reduction, as a human lambda chimeric light chain resulted in normal IgY levels. The reduced level of kappa light chain that is available for secretion may restrict the level of IgY, but that level is sufficient for IgM and IgA, which are normally found at lower levels than IgY. The wild type chicken light chain V gene is more homologous to human lambda (65% amino acid identity to VL3-19 used here) than to human kappa (45% identity to human VK3-15 used here). The human lambda V region may thus be a better match with the chicken CL to fold properly into a functional light chain than the kappa V region. Kappa was initially chosen since the chicken CL to fold properly into a functional light chain than the kappa V region. Kappa was initially chosen since most therapeutic antibodies contain a kappa light chain,\(^2,3\) and it was fused to the chicken CL region to ensure pairing with the chicken heavy chain CH1 domains. The lambda birds were more recently obtained and are under evaluation. Despite the reduction in bulk IgY levels, when compared to WT birds immunized with the same antigen, the OmniChickens were able to produce similar antigen-specific titers and antibodies with similar epitope coverage.\(^8\)

In conclusion, we demonstrated that the active chicken V gene repertoire can be fully replaced by sequences derived from human V genes. The transgenes are expressed in the context of chicken C regions, and are diversified in developing B cells by the native chicken process that includes both gene conversion and somatic hypermutation. The resulting transgenic chickens are phenotypically normal in B cell development and deliver affinity-matured human-sequence antibodies, yet share the immune recognition characteristics of WT chickens, including responsiveness to mammalian-conserved targets and a propensity to generate pan-mammalian cross-reactive antibodies. Interestingly, the “chicken-like” immune recognition is preserved in the OmniChicken despite it being genetically constrained to generate its repertoire based upon a restricted set of V gene frameworks. By focusing on selected frameworks chosen for developability, it is expected that the OmniChicken will yield high quality therapeutic leads at an improved frequency relative to other antibody discovery approaches.

**Materials and methods**

** Constructs used to make OmniChickens**

Production of the IgL and IgH knockouts was previously described.\(^2,23\)

The human kappa construct, SynVK-CK, was identical to the SynVK-C construct previously described\(^25\) except for a modified Kozak consensus sequence in the functional V. The functional V region consists of a rearranged human VK3-15/JK4 gene and was obtained from screening a small library of VK3 family sequences from human PBMC RNA. For the VK pseudogenes, a diverse set of CDRs was selected from the naturally-occurring somatic human sequences found in the NCBI EST database, which was queried with the VK3-15 germline gene. Individual pseudogenes may consist of CDRs from different ESTs, and some framework diversity was also included in some of the pseudogenes. Framework 4 sequence was not included in the pseudogenes. For the human lambda construct, the functional V was swapped for a rearranged VL1-44 gene and VL-based pseudogenes. For the human heavy chain, the functional VH was obtained from screening a small library of VH3 family sequences from human PBMC RNA. The selected V region is based on germline VH3-23 with a few somatic changes and is a rearranged V region consisting of V, D and JH4 elements. For the VH pseudogenes, two versions were made, and OmniChickens contained either one or the other. In construct SynVH-C (used in the BDNF and PGRN campaigns), a diverse set of CDRs was selected from the NCBI EST database, which was queried with the VH3-23 germline gene (Supplementary Fig. 2). Pseudogenes may have CDRs from different ESTs, some framework sequences were included, but no framework 4 sequence was present in the pseudogenes. In construct SynVH-A7 (used in the cocktail immunization), the CDRs were composed of S, Y and W residues, with no changes in the frameworks relative to the functional VH (Supplementary Fig. 2). The pseudogenes were synthesized (Geneziz) with flanking 50 base pair spacers derived from the chicken heavy chain pseudogene locus and cloned into an array, all in the opposite orientation to the functional V. The IgH promoter region was amplified from chicken genomic DNA. The functional V and a short segment of the chicken J-C\(\mu\) intron were synthesized. These parts were assembled with a \(\beta\)-actin promoter and attB site for integration into the attP-neo present in the IgH KO.\(^2,22\)

**Germ cell culture**

Germ cell derivation and culture were performed as previously described.\(^21,38,39\) Re-derived Knockout cell lines IgL KO 229–92\(^23\) and IgH KO 472–138\(^40\) were transfected with the SynVK and SynVH constructs, respectively, using phiC31 integrase as described.\(^34\)

**Breeding**

Animal experiments were done in accordance to Institutional Animal Care and Use Committee (IACUC)-approved protocols and under supervision of the IACUC. To obtain OmniChickens with the genotype SynVK/IgL KO; SynVH/IgH KO, crosses between SynVK/+; IgH KO/+ and IgL KO/+; SynVH/+ birds were
performed. Progeny were genotyped by PCR using DNA obtained from comb biopsy. Males and females were kept for analysis.

**Flow cytometry**

PBMCs were isolated using Histopaque (Sigma, 10771), labeled on ice for 1 h with the following reagents from Southern Biotech: ms anti-Bu1 (8395–01), ms anti-ch IgM (8310–01), ms anti-ch IgL (8340–01), ms anti-ch TCR-γ (8230–01), ms anti-TCRαβ/Vβ1 (8240–01) or ms anti-TCRαβ/Vβ2 (8250–01). All antibodies were diluted in 1% bovine serum albumin (BSA)-phosphate-buffered saline (PBS). Following incubation in primary antibody, samples were washed and then incubated with AF-647 anti-ms IgG (Thermo Fisher, 115-605-205). For double-labeling experiments, direct conjugates (Southern Biotech) were used. Data was collected using an Attune Acoustic Focusing Cytometer (Thermo Fisher) and analyzed using FloJo.

**Elisa**

ELISA plates were coated with ms anti-ch IgA (Southern Biotech, 2330–01), ms anti-ch IgM (Southern Biotech, 8310–01)) or rb anti-ch IgY (Sigma, C2288) at 2 ug/mL in PBS, washed in PBS/0.5% Tween-20 (PBST), and blocked for 1 h in PBST/3% nonfat dry milk. The blocking buffer was aspirated and serial dilutions of plasma in blocking buffer added for 1 h at RT. Plates were washed, incubated with goat anti-ch IgA-HRP (Bethyl, A30-103P), anti-ch IgM-HRP (Bethyl, A30-102P) or anti-ch IgY-HRP (Sigma, A9046) for 1 h at RT, washed, developed with TMB (Thermo Fisher) for 10 min, and the reaction stopped with 1 N HCl. Plates were read at 450 nm on a BioTek microplate reader. Purified IgM (Rockland, 003–0107) and IgY (GenScript, A01010) were used to quantitate plasma Ig titers. IgA was quantified against IgA purified and quantified from chicken plasma (Kaspers Lab, University of Munich).

To assess antigen-specific titer, ELISA plates were coated with 2 ug/mL antigen in PBS. Plates were blocked as described above, incubated with plasma at the indicated dilutions for 1 h, washed, incubated with rb anti-ch IgY-HRP (Sigma, A9046) for 1 h washed, and developed as above.

**Protein and DNA immunizations**

Immunogens were: recombinant human PGRN (Sino Biologicals, 10826-H08H), human BDNF (Sino Biologicals, 50240-MNAS), bovine thrombin (Sigma, T4648), murine IL-13 (Peprotech, 210-10826-H08H), human BDNF (Sino Biologicals, 50240-MNAS), Immunogens were: recombinant human PGRN (Sino Biologicals, 10826-H08H), human BDNF (Sino Biologicals, 50240-MNAS), bovine thrombin (Sigma, T4648), murine IL-13 (Peprotech, 210-10826-H08H), human BDNF (Sino Biologicals, 50240-MNAS), mouse TCR-αβ/Vα2 (8240–01) or mouse TCRαβ/Vβ2 (8250–01). All antibodies were diluted in 1% bovine serum albumin (BSA)-phosphate-buffered saline (PBS). Following incubation in primary antibody, samples were washed and then incubated with AF-647 anti-ms IgG (Thermo Fisher, 115-605-205). For double-labeling experiments, direct conjugates (Southern Biotech) were used. Data was collected using an Attune Acoustic Focusing Cytometer (Thermo Fisher) and analyzed using FloJo.

**Single B cell GEM screening**

We employed a single lymphocyte screening method, the Gel-Encapsulated Microenvironment (GEM) assay (US Patents 8030095 and 84151738) to identify antigen-specific mAbs from immunized chickens. Five μm aldehyde-latex beads (Thermo Fisher) were coated with either purified antigen, or with streptavidin followed by biotinylated antigen, overnight, blocked with 3% milk-PBS, and tested by labeling with plasma from immunized animals. GEMs were prepared containing a single secreting B cell and antigen-coated beads and incubated for 3 h at 37°C in RPMI/10% fetal calf serum containing 2 μg/mL Alexa Fluor 594 anti-ch IgY.

**Single cell PCR to amplify VH and VK regions and clone as scFv-Fc**

Cells secreting antigen-specific mAbs were captured and their VH and VK regions were amplified by a two-step, semi-nested strategy as described. Primer sequences are given in Supplementary Table 1. The variable regions were assembled with human Fc by overlap extension PCR.

**Expression of scFv-Fc antibodies**

Recombinant scFv-Fc was expressed in Expi293 cells as described. Recombinant PGRN was expressed using the same transfection method in 30 mL with 30 μg of DNA.

**Biosensor analysis**

Interaction analysis studies were performed by Array SPRi at 25°C in a running buffer of PBS + 0.05% Tween20 + 1 g/L BSA (unless stated otherwise) using a MX96 SPR imager (IBIS, Netherlands) equipped with Xantec CMD 200M sensor chips. Epitope binning was performed as described previously with the following modifications. All HuMabs were used directly from crude supernatant and typically diluted 10-fold into 10 mM sodium acetate pH 4.5 ± 0.01% Tween-20 (coupling buffer) to amine-couple them onto the chip. The BDNF HuMabs were analyzed using a “premix” assay format in a running buffer of 10 mM Hepes pH 7.4, 150 mM ammonium sulfate, 0.05% Tween-20, and 1 g/L BSA. Briefly, this involved premixing 43 nM binding sites of BDNF with titrating levels of
each HuMab (representing final dilutions of 1/15, 1/30, 1/60, and 1/120 of the supernatant) and injecting these equilibrated mixtures over the coupled HuMab array. Binding responses were compared to those for BDNF alone to determine whether the premixed HuMabs blocked the coupled HuMabs, which were regenerated with a cocktail of 4:1 v/v Pierce Elution buffer (pH2.8) + 5 M NaCl. The bovine thrombin and PGRN HuMabs were analyzed using a “classical sandwich” assay format in a running buffer of PBS pH7.4, 0.05% Tween-20, and 1 g/L BSA and the arrayed HuMabs were regenerated with 75 mM phosphoric acid. By merging the HuMabs with a panel of purified mAb standards from the WT chicken, we inferred the subdomain (or “granulin”) specificities of the PGRN HuMabs by their cross-blockade of the standards. As a complementary approach, the subdomain assignments of the human-specific PGRN HuMabs were determined using a chimeric swap epitope mapping strategy by injecting the HuMabs as analytes over spots coupled with human PGRN, mouse PGRN, five previously described human/mouse chimeras, and a 19-kDa fragment of human PGRN expressing only the D+E subdomains. Epitope binning and mapping of the PGRN HuMabs were performed in the same experiment on the same chip onto which the mAbs and antigen variants were coupled. Binning and mapping data were processed (calibrated, locally-referenced, aligned, and zeroed) in IBIS’s SPRINT software and analyzed in the Epitope Binning Tool v 2.0 (Carterra, Inc., Salt Lake City, UT) to generate network blocking plots.

Affinities of the PGRN HuMabs and WT standards were determined using a capture-based kinetic analysis on an amine-coupled “lawn” of the capture reagent (goat anti-human Fc polyclonal, Southern Biotech, 2047–01). Coupling was performed in the MX96 SPR imager using coupling buffer (as above, pH 4.5) as the running buffer. Three solutions were injected for 7 min, one after another as follows: 1) the surface was activated with a freshly prepared mixture of 40 mM EDC + 10 mM sulfo-NHS (representing a 10-fold dilution of their stock concentrations), 2) the capture reagent was coupled, and 3) excess reactive esters on the surface were blocked with 1 M ethanolamine.HCl pH 8.5, resulting in a final coupling level of about 12,000 response units (or 1,200 millidegrees). The anti-human-Fc-coated chip was then inserted into a continuous flow microspotter (CFM) (Carterra, Inc) and the HuMabs (diluted 3- to 10-fold in PBS + 0.05% Tween-20 + 1 g/L BSA) were captured for 15 min using a 48-channel printhead. The purified WT mAb standards were diluted to 2–5 μg/mL and printed in parallel. Two consecutive prints were used to array 96 mAbs per chip. The printed chip was then redocked into the MX96 SPR imager and primed with PBS + 0.05% Tween-20 + 1 g/L BSA as running buffer. PGRN was prepared as a 3-fold dilution series at nominal concentrations of 0, 0, 7, 22, 67, and 200 nM and injected in increasing concentrations for 3 min each, allowing a 15-min dissociation phase, without any regeneration. Data were processed in SPRINT (as above) with a double-referencing step and the double-referenced data were analyzed globally in Carterra’s kinetic tool “KIT” using a simple Langmuir model with mass transport, allowing each spot its own R_max value. The affinity or equilibrium dissociation constant (K_D) for PGRN interacting with each captured mAb, was determined by the ratio of the kinetic rate constants, K_D = k_d/k_a.

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