

Research Article

Non-Invasive Imaging of Gene Expression and Protein Secretion Dynamics in Living Mice: Identification of Ectopic Prothrombin Expression as Driver of Thrombosis in Cancer

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Abstract

The topology of gene expression and protein localization is a crucial characteristic of life, where the spatiotemporal dynamic of secretory

proteins instruct higher order organization, including the orchestration of developmental and adaptive programs. However tools to non-invasively interrogate the fate of secretory proteins in vivo are scarce. Here we introduce a genetic tagging strategy for in vivo imaging of the secretion and expression dynamics of secretory proteins in living animals. Applying this to a prototypical liver-derived secretory protein, we demonstrate that this approach, combined with optical in-vivo imaging, uncovers extrahepatic prothrombin expression in multiple novel anatomical sites (including testes, placenta, brain, kidney, heart and lymphatic system) and in emerging tumors, resulting in significant amounts of tumor-derived prothrombin in the blood with procoagulant properties. Syngeneic cell lines from this mouse model enable unravelling regulatory mechanisms in high resolution, and in a scalable format ex vivo. Beyond discovering new functions of proteins in a targeted manner, this model allows identifying rheostats in the cross-talk between gene expression and availability of a secretory protein. It is also a valuable resource for uncovering novel (tissue-specific) therapeutic vulnerabilities.

Introduction

Cells synthesize thousands of proteins with diverse functions, which need to be directed to specific locations in accurate amounts at precise time [1]. Nearly one-third of the human proteome is targeted to secretory environments consisting of the endoplasmic reticulum, lysosome, plasma membrane or the extracellular space [2], by a well-operating cargo system [3-4]. Almost all cytokines, hormones, receptors, peptidases, channels, extracellular matrix components, transport proteins and coagulation factors are clients of this machinery [5]. Secretory proteins also possess a central role in anatomical and functional compartmentalization, for example by controlling the function, volume and integrity of the vascular compartment [6-8]. Dysfunction of the production and proper delivery of secretory proteins is the cause of a variety of diseases, including neurodegenerative, developmental and cardiovascular disorders, and perturbations of the hemostatic system [9-14].

In most cells, secreted proteins account for 10-20% of the transcriptome [2]. In contrast, about 40% of all transcripts in the liver encode secreted proteins, reflecting the particular functional demands of this metabolic organ. The liver synthesizes and releases into the bloodstream numerous secretory proteins, including albumin, the most abundant serum protein regulating the colloid osmotic pressure and serving as carrier protein, and most coagulation factors [15]. Accordingly, the impairment of hepatic protein production and secretion results in detrimental consequences [16]. This includes bleeding disorders and venous thromboembolism, a global cause of mortality [17], exemplifying the importance of a well-balanced production of liver-derived secretory proteins [14]. Regulatory mechanisms have evolved to ensure the required functional levels of secretory proteins [18] and to adapt production to conditions of increased turnover and demand [19]. This likely also involves hitherto largely undefined (auto) regulatory mechanisms reminiscent of hormone feedback circuits. Depending on spatial and temporal availability, secretory

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exchange vector harboring an inducible H1 promoter (H1tetO)-driven shRNA cassette along with a genetic element for the constitutive expression of the codon-optimized tetracycline repressor protein (iTetR), and a neomycin resistance cassette was transfected into C57BL/6 ES cell line equipped with RMCE docking sites in the ROSA26 locus. Recombinant clones were isolated using neomycin resistance selection and positive clones harbouring six different shRNAs targeting F2 were pretested for knockdown potency in ES cells (by qPCR analysis). The clone with highest knock-down efficiency was used for the generation of the mouse line. All animal experiments were approved by local authorities, and animals' care was in accordance with institutional guidelines.

Isolation of primary hepatocytes and primary hepatocyte culture

Mice were anesthetized with Ketamine (50 µg/ml, Inresa Arzneimittel GmbH) and the abdomen was opened. Subsequently the portal vein was cannulated to perfuse the liver with 100 ml of prewarmed (37°C) primary hepatocyte isolation (PHI) buffer I (137 mM NaCl, 52 mM KCl, 10 mM HEPES, pH7.4). Upon successful flushing, 50 ml of PHI-buffer II (58 mM NaCl, 52 mM KCl, 1.6 mM CaCl₂, 10mM HEPES, 0.05% collagenase, pH7.6) was used for digestion of the liver. Afterwards, the perfused liver was explanted and stored in PHI-buffer III (137 mM NaCl, 52 mM KCl, 1.6 mM CaCl₂, 10 mM HEPES, pH7.6). The liver was then mechanically dissociated under sterile conditions and dissociated through a 40 µm cell strainer (Corning) and centrifuged (90 g, 3 minutes without break). Next the supernatant was discarded and the liver cells resuspended in 15 ml prewarmed PBS followed by a second round of centrifugation (300 g, 3 minutes without break). The cells were resuspended in William's E medium (Biochrom). Viability was measured by the trypan blue exclusion test. Hepatocytes were then seeded on gelatin-coated culture plates (0.2% gelatin) for 1 hour at 37°C in William's E medium containing 5 % FBS, 1 µM Dexamethasone in DMSO, 100 units/ml Penicillin, 100 µg/ml Streptomycin, 4 µg/ml Human Recombinant Insulin, 2 mM GlutaMAX, 15 mM HEPES at 37 °C in 5%CO₂. The culture medium was changed every 2-3 days. D-Insight positivity of primary hepatocytes was confirmed by luminometric analysis (see below), by FACS based on iRFP fluorescence in a BD FACS Canto (BD Bioscience) and by an Amnis ImageStream Mk II flow cytometer (Luminex) and analyzed using the IDEAS software (Luminex).

Generation and assessment of MCA-induced tumors

Methylcholanthrene (MCA) induced sarcomas were generated through 0.1mg MCA suspension (Sigma #213942) in 30µL corn oil (Sigma #C8267) intramuscularly injected into the left gluteal muscle of D-Insight animals (8-12 weeks old).

Generation of tumor cell lines

Tumors were extracted before they reached diameters >2cm. The tumors were washed in PBS and cut into several fragments and transferred into a 6-Well-Plate containing digestion medium (RPMI, 20 % FBS, 100 units/ml Penicillin, 100 µg/ml Streptomycin, 8 mg/ml Collagenase) and incubated for 24 hours at 37 °C in 5%CO₂.umor fragments were mechanically dissociated by pipetting up and down, and then filtered through 40 µm cell strainers. The suspension was centrifuged at 1000 x g for 5 minutes and the obtained cell pellet was resuspended in DMEM (Biochrom). Viability was measured by the trypan blue exclusion assay. Fibrosarcoma cells were seeded on

culture plates in DMEM containing 20 % FBS, 3.7 g/l NaHCO₃ 4.5 g/l D-Glucose at 37 °C in 5%CO₂. The medium was changed every 48 h.

Generation of primary fibroblast cultures

Mice ears were washed in PBS and cut into several fragments. The fragments were transferred into a 6-well-plate containing digestion medium (RPMI, 20 % FBS, 100 units/ml Penicillin, 100 µg/ml Streptomycin, 8 mg/ml Collagenase) and incubated for 24 hours. Ear fragments were mechanically dissociated by pipetting up and down and filtered through 40 µm cell strainers. The suspension was centrifuged at 1000 x g for 5 minutes and the obtained cell pellet was resuspended in DMEM. Viability was measured by the trypan blue exclusion assay. Fibroblasts were seeded on culture plates in DMEM containing 20% FBS, 3.7 g/l NaHCO₃, 4.5 g/l D-Glucose at 37 °C in 5%CO₂. The medium was changed every 48 h.

In vivo Imaging

Non-invasive in vivo imaging of D-Insight mice was performed under Ke/Xy narcosis (intraperitoneal (i.p.) injection of ketamine (Ke, 80-120 mg/kg body weight) / xylazine (Xy, 5-10 mg/kg body weight) cocktail) using an IVIS Spectrum In Vivo Imaging System (Perkin Elmer, Rodgau, Germany) with images acquired using the Living Image® software package (Perkin Elmer).

The dorsal or ventral images of mouse bodies, ventral images of mice in operation, ex vivo imaging of the same mice (after euthanasia by overdose application of Ke/Xy narcosis) or postmortem imaging of certain organs were performed 30 min after i.p. injection of 100 µl ready-to-use coelenterazine (RediJet Coelenterazine H Bioluminescent Substrate, PerkinElmer, Rodgau, Germany) or 10 min after i.p. injection of 100 µl ready-to-use luciferin (RediJet D-Luciferin Bioluminescent Substrate, PerkinElmer, Rodgau, Germany) respectively. The pseudo color luminescent images (blue, green, yellow, and red from least to most intense) were overlaid on the grayscale photographic images. Quantification of luminescence was determined by calculating the Average Counts as Total Counts/Number of pixels for a defined region of interest.

Luminometry Assays

100000 cells or 200 mg tissue were lysed in 100-500 µl Passive Lysis Buffer (Promega) for 15 min at room temperature on an orbital shaker. 50 µl of the lysate, plasma or media samples, were assayed in white 96 well plates in triplicate by the addition of 50 µl Bright-Glo™ Firefly luciferase assay system (Promega) or freshly prepared 40 ng/ml Coelenterazine in Renilla-Glo buffer (Promega). Alternatively 20 µl samples were assayed using a Dual-Glo Luciferase Assay System using manufacturers protocol (Promega). Luminescence (RLU) was measured immediately using a GloMax® Discover Microplate Reader (Promega).

Protein Transport Inhibitor Treatment

200000 cells were seed in Hepatocytes-Williams E medium (Gibco, 10043282) containing 10% FCS, 100 units/ml Penicillin and 100 µg/ml Streptomycin and incubated at 37 °C in 5% CO₂. After 3 h, 0-0.5 µg/ml Brefeldin A was added to the medium and cells were further incubated for 2 days. Luminometry assays were then performed as described above.

Image Stream

For real time imaging of iRFP signal in DI hepatocytes and fibrosarcoma cells image stream was used. Cells were trypsinized and diluted in FACS buffer (PBS (1x), Bovine serum albumin (BSA, 0.5%), fetal bovine serum (FBS, 0.1%), Sodium azide (NaN₃, 0.1%)) to a final concentration of 1 million cells per 500 μ L. B6 hepatocytes and fibrosarcoma cells were used as controls. 1 μ L of DAPI (1:10000 diluted in PBS) was added to 20 μ L cell suspension before loading samples on the ImageStreamX MkII imaging cytometer. Single cell population and live cells were gated by considering the signal generated in DAPI 405 channel. For activation of iRFP signal in fibrosarcoma cells, 3-5 μ L of biliverdin hydrochloride (chromophore for iRFP signal activation, final concentration 0.002%, Sigma; 30891) added to cells and incubated for 10 mins at 37 °C before imaging. As liver cells are known to produce sufficient amounts of bilirubin no additional biliverdin was used for iRFP activation in hepatocytes population. Living cells were sorted for iRFP positive signal and were quantified by gating the cells based on signal from 702/86 channel. The INSPIRE software was used to analyze the data and graph prism was used to plot the graphs.

LPS-treatment

Mice were injected intra-peritoneal with 1.0 to 1.5 mg/kg LPS (Sigma Aldrich). After 3 to 48 h mice were sacrificed by isoflurane inhalation and liver and plasma samples obtained. Samples were assayed by luminometry as described above.

F2 qRT-PCR

Total RNA recovered from cells using peqGOLD TriFast reagent (Peqlab) and cDNA produced using the RevertAid Reverse Transcriptase kit (ThermoFisher Scientific) as per manufacturer's instructions. qRT-PCR was performed using the Blue S'Green qPCR Kit (Biozym) with primers for F2 (GGTGAACCTGCCATTGTAGA, TCCTC-GCTTGGTGTCAATTCA) and the housekeeper gene GAPDH (AG-GTCGGTGTGAACGGATTG, TGTAGACCATGTAGTTGAG-GTCA) as per manufacturer's instructions. Cycling (95°C for 2 min; 95°C for 30 sec; 60°C for 30 sec; 72°C for 30 sec and followed by a melting curve) was performed in a CFX Real-Time Detection System (Bio-Rad) and relative expression calculated using the CFX Manager Software (Bio-Rad).

F2 RNA in situ hybridization (FISH)

Cryosections were generated from murine organs and tissues. Briefly, mice were sacrificed by isoflurane inhalation and organs were washed in PBS before placing in Tissue-Tek® Cryomolds (25×20×5mm or 15×10×5mm; Sakura). The specimens were immediately covered with Tissue-Tek® O.C.T. compound, put on dry ice until frozen and stored at -80°C until sectioning. Cryosections (12 μ m) were produced using a cryostat, mounted onto Superfrost®plus slides (Thermo Fisher Scientific) and fixed overnight at 4°C in 4% formaldehyde (FA).

The Thermo Fisher Scientific ViewRNA ISH Tissue 1-Plex Assay was used to visualize the F2 mRNA expression in cryosections. Briefly, slides were washed twice in PBS and dehydrated in increasing EtOH concentrations (50%, 70%, 100%; 10 min each). After drying, a hydrophobic barrier was drawn around the specimen using the ImmEdge pen (Vector). Slides were boiled (85-95°C) for 30-45s in pretreatment solution and washed twice with PBS, followed by

incubation in 0.2M HCL (RT for 10 min). After washing with PBS, slides were incubated (10 min, RT) with Protease solution (diluted 1/100 in prewarmed PBS), followed by rinsing in PBS and 3 minutes incubation in 4% FA. Thereafter, all following hybridization and amplification steps were carried out at 40°C in a humidified chamber.

After washing twice with PBS and once with ddH₂O, the hybridization of the probe set was carried out by incubation with the probe set working solution (probe set diluted 1/30 with prewarmed Probe set diluent) for 2.5-3h at 40°C. After hybridization, slides were washed 3x in washing buffer for 2 min at room temperature and under vigorous shaking. The slides were then incubated for 40 min in Pre-Amp solution diluted 1/100 in Amplifier Diluent, followed by a second washing step as indicated above. The slides were then incubated for 40 min in Amp solution diluted 1/100 in Amplifier Diluent. After a third washing step, the slides were incubated in Label Probe-AP diluted 1/500 in Label Probe Diluent for 40 min, followed by washing 3x in washing buffer for 3 min each with continuous shaking. AP-Enhancer solution was added to the samples and incubated for 5 min at room temperature. The slides were then incubated with freshly prepared detection substrate, consisting of half a tablet of Fast Red Substrate resuspended in 2.5 ml Naphthol buffer, for 30-40 min at 40°C in the humidified chamber. Slides were then washed 2x in PBS and kept in 4% FA for 5 min. After washing in ddH₂O, counterstaining in haematoxylin for 10-30 sec and destaining in ddH₂O, slides were dried 20 min at room temperature in dark environment, mounted with Fluoromount-G with DAPI (Invitrogen), covered with coverslips and dried again for 15 min. Nail polish was applied to seal the edges and slides were stored at 4°C covered with aluminum foil.

F2 activity, APTT measurements, Clinical Chemistry

F2 activity of tumor-bearing and tumor-free animals was measured with ACL Top instruments and a KC4™ Coagulation Analyser. Analyses of transaminases (GOT/GPT) have been carried out on an Abbott Alinity C analyzer.

Diphtheria-Toxin treatment

The dose of Diphtheria toxin (DTX) was titrated in the range 0-4 ng/g body weight per animal with 2 ng/g being defined as optimal and used for the shown experiment (injected i.p.).

Epigenetic modulator treatment of D-Insight fibrosarcoma cells 100000 D-Insight fibrosarcoma cells were seeded in 6 Well plates in DMEM containing 10% FBS, 3.7 g/l NaHCO₃, 4.5 g/l D-Glucose and incubated at 37 °C in 5% CO₂ for 24 hours. Media was then changed to include an epigenetic modulator (either 10 nM (+)JQ1, 10 μ M Azacitidine, 10 μ M Decitabine, 10 μ M Lomeguatrib, 10 nM Panobinostat, 10 nM Quisinostat, 10 μ M RG108 or 10 μ M Zebularine in DMSO) and incubated for 2 - 7 days and luminometry performed as described above.

Statistical Analysis

All statistical analyses and production of graphs were performed using Prism 9 (GraphPad Software).

Results

Thrombin is the key serine protease involved in blood coagulation and hemostasis. It is synthesized in the liver as an inactive 72-kDa precursor protein (prothrombin) and secreted into the blood circulation. Upon activation, thrombin converts fibrinogen into fibrin, the

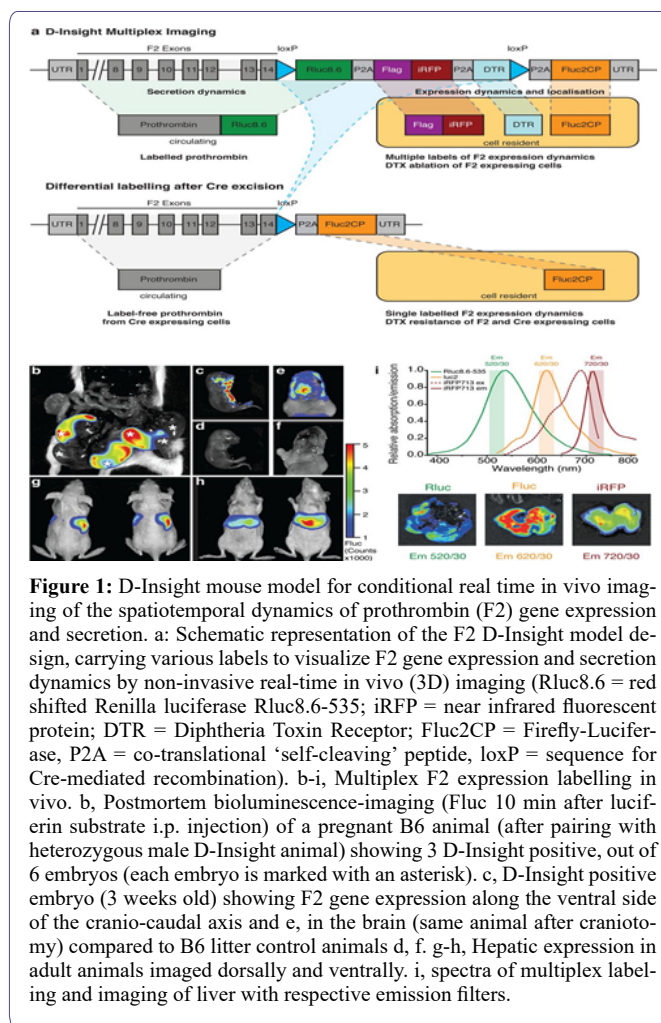
main component of blood clots, and activates platelets. Besides its role in hemostasis [34], thrombin is involved in numerous other processes, including embryonic development, angiogenesis, wound healing, inflammation, atherosclerosis and tumor biology [35-36]. This reflects the broad action of thrombin on membrane-bound G-protein coupled Protease Activated Receptors (PARs) [37]. However, many of these actions naturally affect cells residing in the extravascular compartment [34]. This questions the prevailing observation of prothrombin synthesis being confined to the liver [2].

Here we set out to generate a multimodal imaging reporter animal to characterize the spatiotemporal dynamics of this pleiotropic liver-derived protein in vivo. Specifically, our strategy aimed at creating a reporter for (1) identifying and quantifying (non) canonical origins of prothrombin expression, and (2) disentangling gene expression and protein secretion dynamics under control of endogenous regulatory mechanisms. Further it ought to feature a (3) differential labeling of the secretory protein (depending on the origin where it is expressed and secreted), provide a (4) component to explore the downstream function(s) and serve as (5) resource for isogenic primary reporter cell lines for further in-depth studies, including regulatory pathways by high content analyses in a scalable format *ex vivo*.

To generate this versatile tool, we employed a multicistronic reporter strategy targeting the endogenous prothrombin (F2) gene locus in B6 mice (Figure 1a). We generated a conditional knock-in allele for in-frame insertion of two luminescence reporters, Rluc8.6-535 [38] and Fluc2CP (with the latter harboring two protein destabilization domains, hCL1 and hPEST, for improved reporter dynamics), a bright and stable near-infrared Fluorescent Protein (iRFP) for in vivo imaging [39] and a human Diphtheria-Toxin Receptor (DTR) directly upstream of the translation termination codon and the 3' untranslated region. By using highly efficient viral P2A co-translational 'self-cleaving' peptides [40], the endogenous prothrombin and the inserted reporters are co-expressed as individual proteins in a strict 1:1:1 stoichiometry [41]. While prothrombin is tagged by in-frame fusion to Rluc and secreted into the circulation, the Flag-iRFP, the DTR and the Fluc are cell-resident (and thereby 'label' cells that express prothrombin). For the downstream functional exploration of prothrombin and/or the cell(s) expressing this secretory protein, a BRET-compatible Rluc label was chosen (to allow for ligand-receptor interaction [42] and compartment colocalization studies [43] in addition to the DTR for targeted cell depletion with Diphtheria toxin (DTX) [44]. For a Cre-mediated excision [45], two loxP-sites were inserted upstream of the Rluc and downstream of the DTR. This enables differential labeling of the canonical (liver-derived) and non-canonical (extrahepatic) prothrombin and a conditional DTX-mediated depletion of prothrombin-expressing cells.

Despite the extensive genetic modifications, we obtained viable reporter animals in expected Mendelian ratios (Figure 1b). The obtained Fluc signals are specific for the transgene, and in the developing embryo project to the ventral part of the cranio-caudal axis (Figure 1c, d) and the brain (Figure 1e, f). This corresponds to earlier reports supposing prothrombin expression at various locations in the developing embryo [46] and a function in the nervous system [47]. In contrast, adult animals show strong signals (Fluc, Rluc and iRFP) that co-localize with the liver (Figure 1g-i), corresponding to the hepatic expression of prothrombin in adults (www.proteinatlas.org).

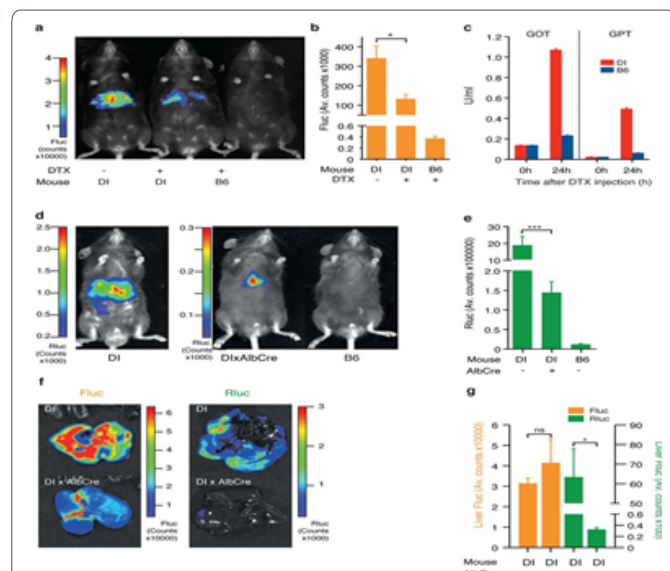
To validate the functionality of the DTR introduced with our multicistronic transgene, we next administered DTX to the reporter and



B6 controls animals. Expectedly, we observe that 24h after DTX injection the Fluc signal declines in the liver (Extended Data Figure 1a,b), reflecting a DTX-mediated disintegration of this organ. This is mirrored by a significant induction of the transaminases (GOT, GPT), reflecting liver tissue damage (Extended Data Figure. 1c). In contrast, B6 animals that do not harbor the human DTR do not show an induction.

For most applications, the DTX-mediated ablation of prothrombin expressing cells (primarily targeting the liver) is incompatible with a functional assessment of minor sites of prothrombin expression. Additionally, we aimed to exploit a differential labeling strategy that allows us to distinguish prothrombin produced in the liver from extrahepatic sites (Figure. 1a). To validate the Cre-mediated conditional labeling, we crossed D-In-sight animals with a liver-specific Cre-expressing line (Albumin-Cre) and analyzed the bioluminescence of the offspring. Expectedly, we obtained animals in which the Rluc fused in-frame to the prothrombin can no longer be detected, while Fluc expression remained intact (Extended Data Figure 1d-g). This manipulation also results in the excision of the DTR (Figure 1a), which enables a selective DTX-mediated ablation of prothrombin-expressing extrahepatic cells. The elimination of Rluc and DTR thereby introduces a functional component to further explore downstream features of extrahepatic prothrombin and/or the function of prothrombin-expressing cells in vivo.

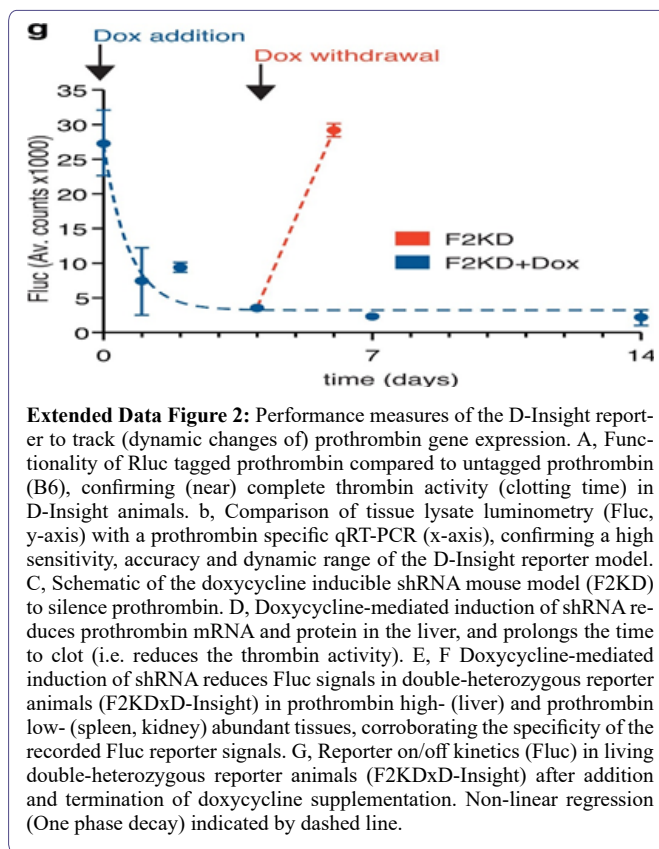
Our reporter is designed to discriminate between prothrombin expression and secretion (Figure 1a). Accordingly, while Rluc and Fluc can be detected in liver lysates, only the Rluc signal can be found in the plasma (Figure 2a). Comparing plasma thrombin activities of B6 and D-Insight reporter animals revealed almost identical levels (Extended Data Figure 2a), demonstrating that the Rluc tag per se does not impede protein transport nor thrombin function.



Extended Data Figure 1: Functional customization of the D-Insight model through Diphtheria toxin receptor mediated targeting and Cre-mediated reporter excision. a-c Validation of Diphtheria Toxin (DTX) mediated targeting of reporter expressing cells (DI +/-DTX), showing reduction of liver Fluc signal b, and a corresponding release of liver derived transaminases (glutamic oxaloacetic transaminase (GOT), glutamic pyruvate transaminase (GPT)) 24 h after DTX injection in comparison to a wildtype B6 animal c. d-g Validation of Cre-mediated excision of the Rluc-DTR reporter cassette with D-Insight animals after pairing with a liver specific Cre-line (AlbCre). This results in a significant reduction of liver-derived Rluc bioluminescence (e, f, g), confirming functionality of the conditional multiplex reporter labeling strategy (Figure 1a).

Next, we aimed to validate the reporter with established modifiers of prothrombin expression. Prothrombin is induced during septicemia by a regulatory mechanism controlling RNA processing [19]. We injected lipopolysaccharides (LPS) into reporter animals and observed a significant upregulation of prothrombin expression (Fluc intensity) in the liver 3 hours after LPS-injection (Fig. 2b) - reminiscent of earlier reported RNA induction. While this induction is rapid, the abundance of prothrombin in plasma (Rluc) increases with retarded kinetics 12 hours after LPS-injection (Figure 2b). This demonstrates that the reporter mouse model recapitulates established prothrombin expression dynamics [19-48]. By differential labeling and quantification of the expression and secretion dynamics, it allows disentangling these processes in vivo.

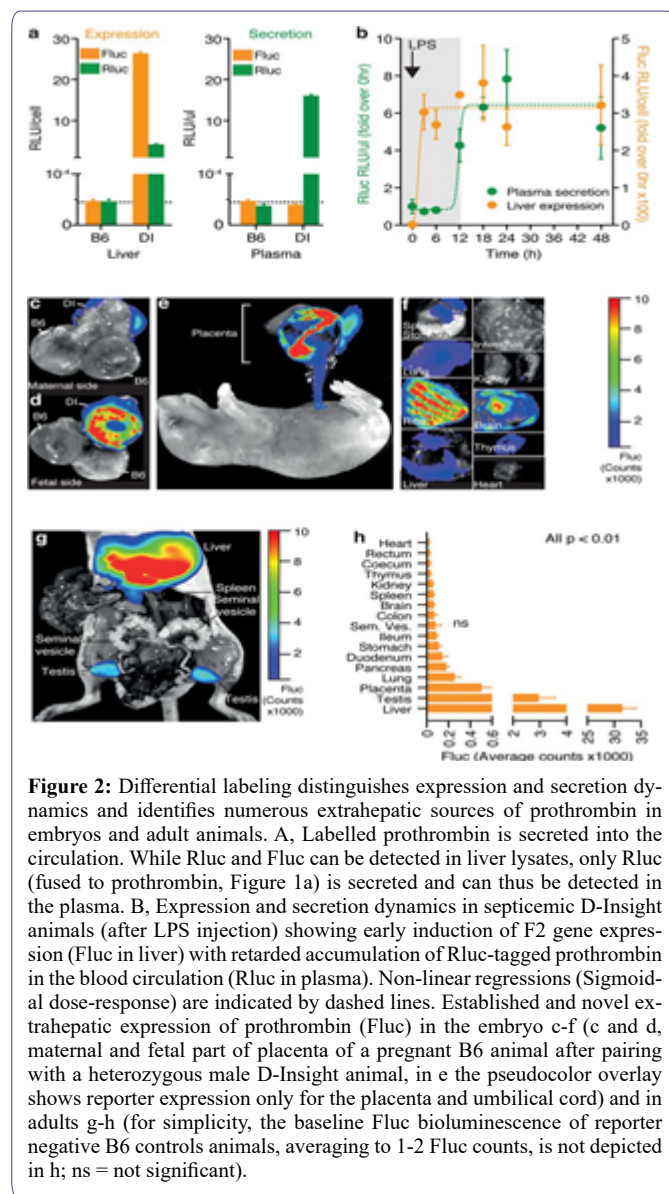
We next tested the accuracy and sensitivity of the reporter system, and whether new anatomical origins of this prototypic liver-derived protein can be detected. The knockout of prothrombin is embryonically lethal with phenotypes associated with impaired vascular integrity during development [49-50]. Based on this, we expected to retrieve prothrombin expression characteristics that can be associated with these findings. We investigated prothrombin expression in developing embryos, and observed that the fetal part of the placenta (and the



Extended Data Figure 2: Performance measures of the D-Insight reporter to track (dynamic changes of) prothrombin gene expression. A, Functionality of Rluc tagged prothrombin compared to untagged prothrombin (B6), confirming (near) complete thrombin activity (clotting time) in D-Insight animals. b, Comparison of tissue lysate luminometry (Fluc, y-axis) with a prothrombin specific qRT-PCR (x-axis), confirming a high sensitivity, accuracy and dynamic range of the D-Insight reporter model. C, Schematic of the doxycycline inducible shRNA mouse model (F2KD) to silence prothrombin. D, Doxycycline-mediated induction of shRNA reduces prothrombin mRNA and protein in the liver, and prolongs the time to clot (i.e. reduces the thrombin activity). E, F Doxycycline-mediated induction of shRNA reduces Fluc signals in double-heterozygous reporter animals (F2KDxD-Insight) in prothrombin high- (liver) and prothrombin low- (spleen, kidney) abundant tissues, corroborating the specificity of the recorded Fluc reporter signals. G, Reporter on/off kinetics (Fluc) in living double-heterozygous reporter animals (F2KDxD-Insight) after addition and termination of doxycycline supplementation. Non-linear regression (One phase decay) indicated by dashed line.

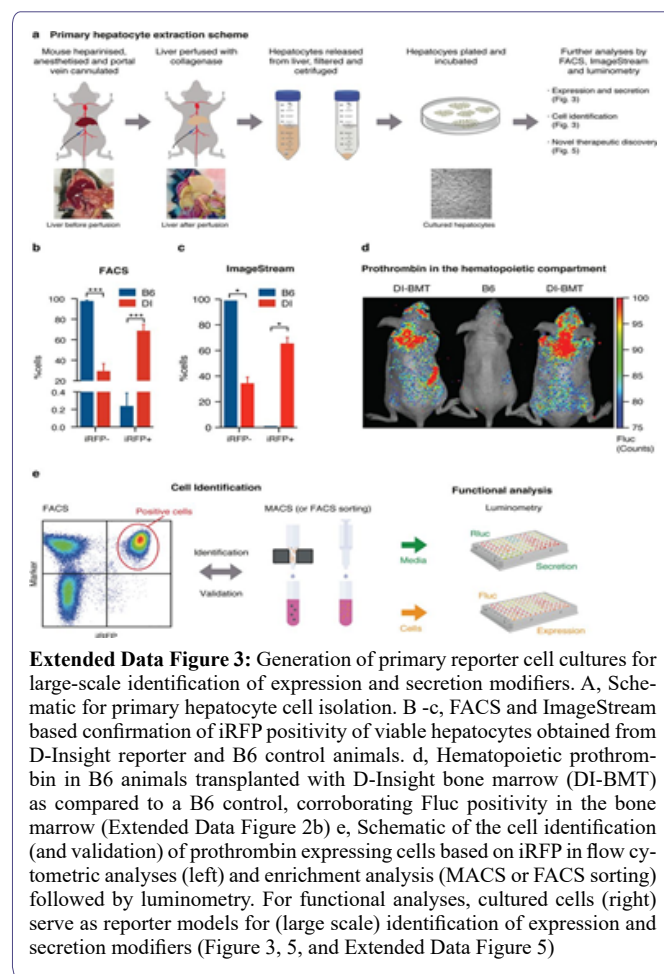
umbilical cord) expresses significant amounts (Figure 2c-e). We also identified expression in a variety of other organs in the developing embryo (Figure 2f). We further extended these studies to adult animals. We identified significant amounts of prothrombin expression in the testis (Figure 2g) and at numerous previously unreported extrahepatic locations (Figure 2h), including the lung, pancreas, lymphatic system (spleen), kidney and heart.

To further confirm the accuracy of the reporter system, we compared Fluc signals of various tissues from D-Insight animals and compared this to prothrombin RNA abundance (Extended Data Figure 2b). This revealed that the reporter activity closely mirrors prothrombin gene expression over more than 2 orders of magnitude, documenting a high dynamic range and sensitivity. To control the specificity (above background) and on/off-kinetics of the reporter, we finally generated a doxycycline inducible short hairpin (sh) RNA mouse model targeting the endogenous prothrombin gene (Extended Data Figure. 2c). Upon doxycycline administration, this model (F2KD) reduces endogenous (pro) thrombin RNA, protein and activity down to 10-25% (Extended Data Figure. 2d). We next crossed this model with the reporter mice and determined luminescence with and without doxycycline administration. This analysis confirms the specificity of the Fluc signal in a prothrombin high abundant (liver) as well as in select low abundant tissues (kidney, spleen, Extended Data Figure 2e, f). Further, the Fluc kinetics in the liver upon doxycycline addition and withdrawal corroborates the rapid on/off-kinetics of the reporter (Extended Data Figure 2g), which is suited for real time imaging of dynamic expression alterations and resulting functional effects (Figure 2b). Altogether this demonstrates the functionality of the multiplex imaging strategy to disentangle the expression and secretion dynamics of a prototypic liver-derived secretory protein in vivo. It also uncovers prothrombin expression in a variety of hitherto unknown extrahepatic locations.



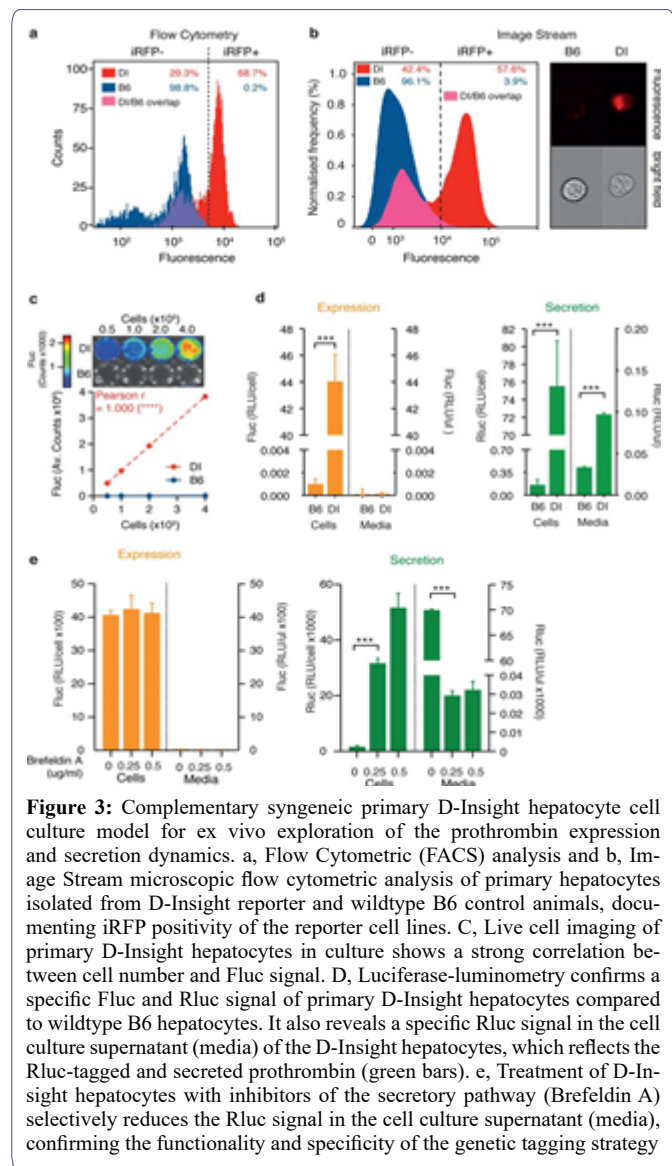
Cell-based reporter systems are potent tools to elucidate disease mechanisms and to identify novel therapeutic vulnerabilities [51]. This becomes extremely efficient if such experimental set-ups can be carried out additively on the same genomic background. To test the in-principle applicability, we generated complementary syngeneic cell lines derived from our reporter animal. We used a modified extraction protocol [52] to isolate primary hepatocytes from D-Insight and B6 control animals (Figure 3, Extended Data Figure 3a). After initial cultivation, we confirmed purity, iRFP and Fluc positivity of reporter hepatocytes using multispectral imaging flow cytometry (Figure. 3a, Extended Data Figure. 3b), Image Stream fluorescence microscopy (Figure. 3b, Extended Data Figure. 3c), luminometry and live cell imaging, where serial dilutions produced Fluc signals correlating with the number of cells (Figure 3c). Recapitulating the findings obtained in vivo (Figure. 2a), we also detected Fluc and RLuc in hepatocyte cell culture lysates (Figure 3d). Conversely, in the media only RLuc can be detected reflecting the secreted RLuc-tagged prothrombin, while Fluc remains cell-resident (Figure 3d). Further, treating these cells with the protein transport inhibitor brefeldin A leads to

a significant reduction of RLuc in the media (Figure 3e), while, expectedly, the RLuc tagged-prothrombin accumulates in the cell. Together this confirms the functionality of this cell-based multimodal reporter imaging approach (Figure 1a) enabling the unraveling of mechanisms governing the expression and secretion dynamics of this liver-derived secretory protein in high resolution and in a scalable format ex vivo. Additionally this multimodal approach provides multiple avenues for identification of novel cellular sources of prothrombin (e.g. flow cytometric analyses for the identification and functional characterization of prothrombin positive cells in the spleen and the lymphatic system (Figure 2h) or hematopoietic (bone marrow) compartment (Extended Data Figure 2b, 3d, e).



We next performed a proof-of-concept experiment to explore how the reporter mouse model performs in detecting potentially new locations of extrahepatic prothrombin expression with pathogenic relevance. Up to 20% of all cancer patients develop venous thromboembolism, reflecting a procoagulant disbalance of the hemostatic system with a significant incidence of fatal outcomes [53-54]. We therefore employed a methylcholanthrene (MCA) tumor induction model to assay whether prothrombin is potentially expressed and secreted by emerging tumors. In MCA challenged reporter animals, we first observed that the emerging tumors express prothrombin (Figure. 4a). We next transplanted these tumors into B6 mice and obtained Fluc signals that increase with time and volume of the transplanted tumor (Figure. 4b, Extended Data Figure 4a, b). This suggests that the initial Fluc signals in the tumor unlikely originate from tumor-associated

macrophages or lymphocytes frequently found in tumors [55]. To consolidate D-InSight-reporter and prothrombin expression, we analyzed tumor cell cultures obtained from these animals (Figure 4c-f). Using multispectral imaging flow cytometry and luminometry, we confirm reporter iRFP positivity (Figure 4c, Extended Data Figure 4c) and Fluc expression (Figure 4d). Using fibrosarcomas of the genetic background in which prothrombin expression can be depleted, we directly confirm endogenous prothrombin mRNA expression by F2 RNA-FISH (Figure 4e, f). Surprisingly, probing further murine and human tumors we also find extrahepatic prothrombin expression in several entities (Extended Data Figure 4d-f), suggesting that this finding may be relevant in tumor biology across species and entities [56].



A, MCA-induced fibrosarcomas in D-InSight mice, showing tumor-associated prothrombin expression (Fluc). B, Tumor-associated prothrombin expression is transplantable and increases with tumor volume. Non-invasive in vivo whole body bioluminescence imaging of Fluc of immunocompetent albino hairless (AH) mice after s.c. transplantation of D-InSight sarcomas at day 10, 20 and 30, respectively (y-axis depicts Fluc, the x-axis displays the tumor volume, n>3 animals per time point). Non-linear regression (Exponential growth

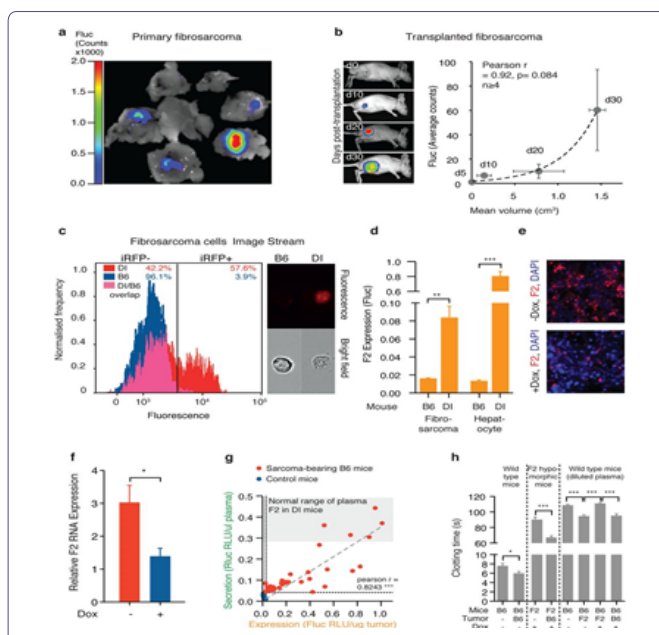
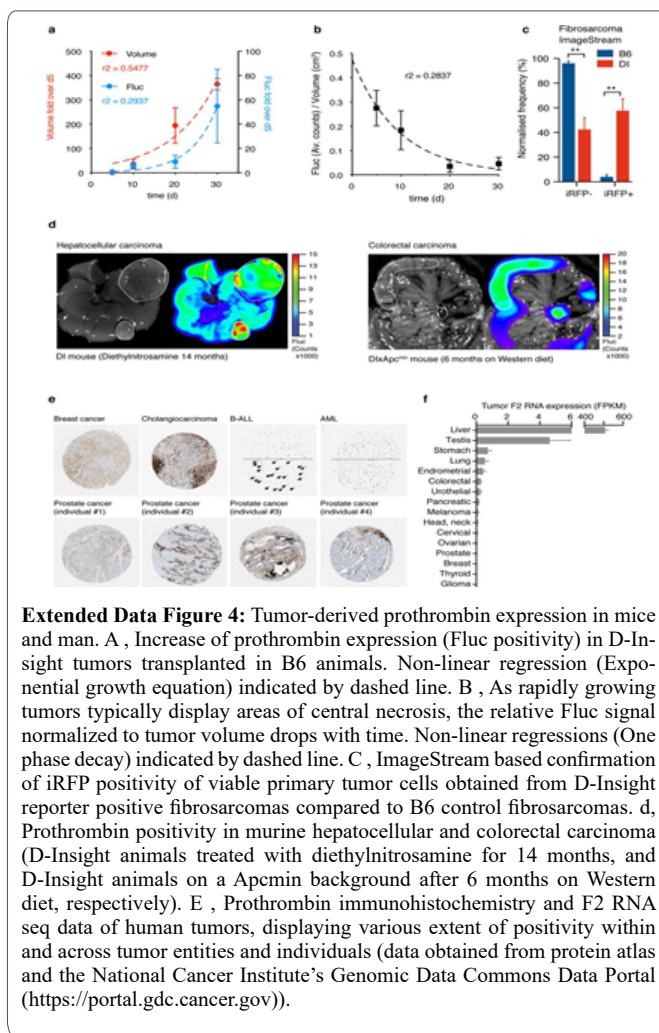


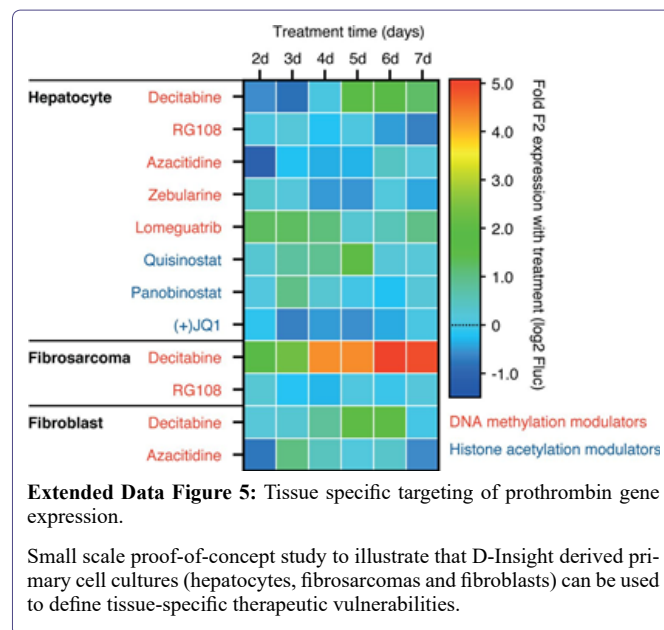
Figure 4: Identification of new locations of extrahepatic prothrombin expression with procoagulant function.

equation) indicated by dashed line. c, d, ImageStream and luminometry of fibrosarcomas confirms iRFP positivity and Fluc expression for D-Insight fibrosarcomas. e,f, Tumor-intrinsic F2 expression confirmed by F2-RNA-FISH with and without doxycycline-mediated depletion of F2. G, Tumor-derived prothrombin is secreted into the blood circulation. The D-Insight reporter Fluc signal (x-axis) of sarcoma bearing mice (red) reflects the amount of prothrombin expressed in tumor cells, which positively correlates with the amount of tumor-derived prothrombin secreted into the circulation (y-axis, Rluc in plasma; signals obtained in muscle tissue and plasma, respectively, for B6 mice (blue) serve as non-specific background control). Notably, D-Insight sarcoma bearing B6 mice (red) express and secrete significant amounts of prothrombin, reaching Rluc plasma levels comparable to liver-derived prothrombin of D-Insight control mice (grey shaded corridor). h, Ectopically expressed prothrombin by the tumor is functionally active (assessed by KC4™ Coagulation Analyser; comparing blood plasma clotting time of F2-KD (F2), B6 mice (B6), F2-KD sarcoma bearing B6 mice and B6-sarcoma bearing F2-KD mice with (+) and without (-) administration of a doxycycline (Dox) rich diet for 3 weeks).

We next explored whether our model system provides further insights substantiating the functional dimension of this unexpected finding. We tested if tumor-derived prothrombin is secreted into the blood circulation. We transplanted D-Insight positive fibrosarcomas into B6 mice and determined Fluc and Rluc levels in the tumor and plasma, respectively. Surprisingly, we observed a strong correlation (Fig. 4g). Comparing these signals with Rluc in tumor-free D-Insight control animals reveals a substantial amount of tumor-derived Rluc that can be detected in the plasma, in concentrations comparable to tumor-free reporter animals (Fig. 4g). This suggests that tumor-derived prothrombin is quantitatively significant; it reaches the intravascular compartment to a similar extent as prothrombin constitutively synthesized in the liver.

The biogenesis of prothrombin involves post-translational gamma-carboxylation in the liver, which is required for full functional activity [57]. To determine the potential functionality of tumor-derived prothrombin, we first assessed whether all enzymes involved in the vitamin K cycle are expressed. We confirmed expression for the gamma carboxylase (GGXC), the NAD(P)H quinone hydroxylase 1 (NQO1), the vitamin K epoxide reductase complex subunit 1 (VKORC1) and the vitamin K epoxide reductase complex subunit 1 like 1 (VKORC1L1) in the tumor at levels similar (or higher) to the liver (not shown). We also applied thrombin activity measurements and analyzed plasma samples obtained from either normal (B6) or prothrombin (F2) hypomorphic animals with and without implanted tumors that express prothrombin (Figure 4f). To that end, we used wildtype B6 and F2 KD fibrosarcomas in which prothrombin expression can be selectively depleted (Figure 4e,f). We identified that tumor-derived prothrombin confers a procoagulant function in B6 mice (Fig. 4h, compare bars 1 and 2). Accordingly, we also observed that tumor-derived prothrombin compensates for low thrombin activity in prothrombin hypomorphic animals (Figure. 4h, bar 3 and 4), and that this compensation is lost, when the prothrombin expression in the tumor was silenced by the shRNA (Fig. 4h, bars 5-8). This suggests that tumor-derived prothrombin is functionally active. This may also perturb the well-balanced equilibrium between pro- and anticoagulatory activities in the plasma [58-59] and could thereby contribute to the hypercoagulable state frequently observed in cancer patients, ultimately leading to detrimental consequences including thromboembolic death [53-54].

Understanding tissue-specific gene regulation [60] is key to tissue-tailored therapies in cancer and beyond [61]. Based on our findings, a therapeutic strategy targeting the expression of such a secretory protein in a tissue-selective manner would be desirable. We therefore finally tested the applicability of this approach and conducted a small-scale chemical screening employing a culture of primary hepatocytes, fibrosarcoma tumor cells and fibroblasts (serving as controls) obtained from the D-Insight reporter animal (Figure 3,4). Using in-plate luminometry in a 96-well format in our screening, we surprisingly identified a DNA methyltransferase inhibitor (Decitabine), which shows a long-lasting and highly selective modulation of prothrombin gene expression in a tissue dependent-manner affecting fibrosarcomas, while leaving the expression in the liver and fibroblasts unaltered (Figure 5, heatmap and Extended Data Figure 5). This points to tissue-selective regulatory mechanisms governing the expression of this secretory protein with a multitude of actions in various systems and disease mechanisms, ranging from atherosclerosis to tumor biology [35-36]. It also illustrates the versatility of this model to dissect disease mechanisms and to uncover novel therapeutic vulnerabilities (Figure 5).



Discussion

The liver is the prime organ producing secretory proteins. While protein targeting is achieved by a well-characterized cargo system 3, 4, new mechanisms of unconventional protein secretion are being discovered [62-63]. Owing to the importance of secretory proteins in a variety of diseases [9-14], understanding regulatory mechanisms on a systems and cellular level may help elucidating underlying biology and pathophysiological processes, including the identification of novel therapeutic avenues [23].

Here we introduce a conditional, multiplex imaging reporter mouse model to disentangle the expression and secretion dynamics of a prototypic liver-derived secretory protein in vivo. Our findings illustrate the versatility of this reporter model for detecting and illuminating processes of this secretory protein in the context of development and various disease mechanisms. They also document the autonomous expression of a secretory protein that becomes harmful if

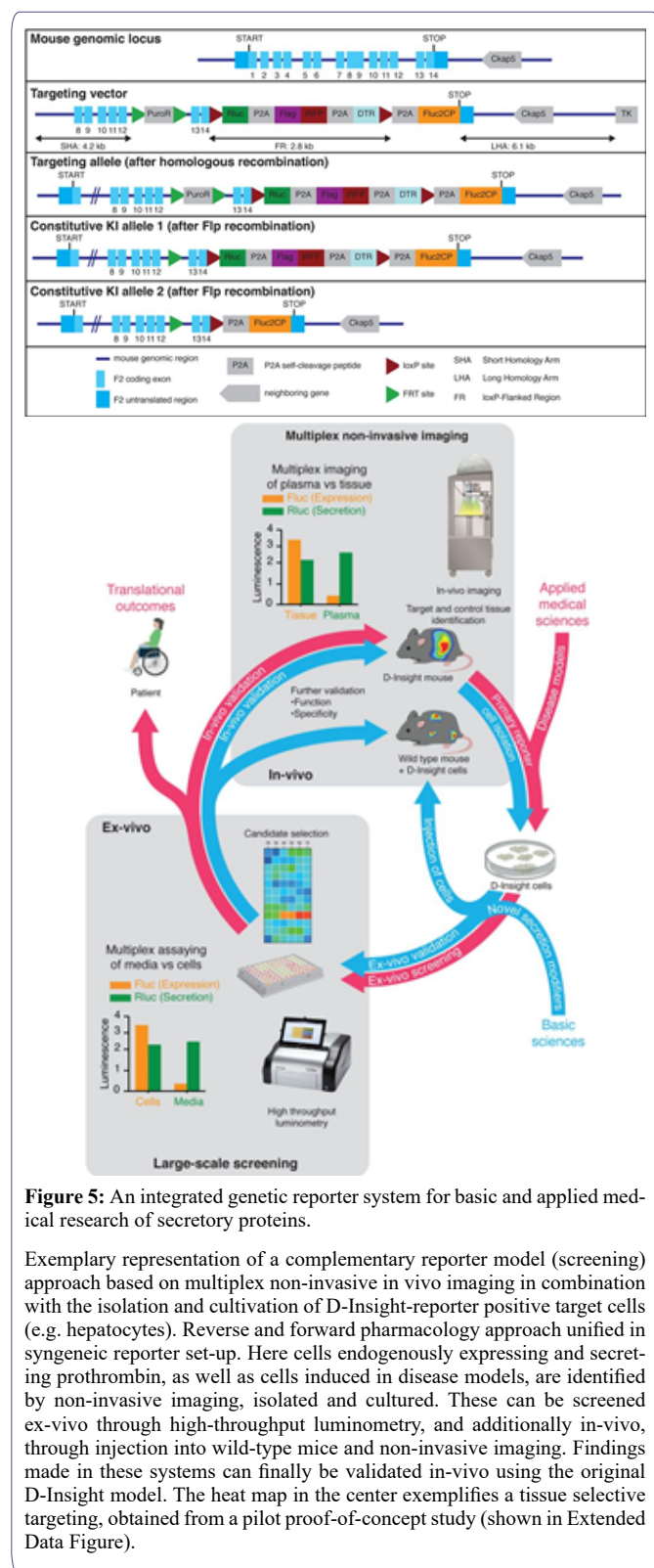


Figure 5: An integrated genetic reporter system for basic and applied medical research of secretory proteins.

Exemplary representation of a complementary reporter model (screening) approach based on multiplex non-invasive in vivo imaging in combination with the isolation and cultivation of D-Insight-reporter positive target cells (e.g. hepatocytes). Reverse and forward pharmacology approach unified in syngeneic reporter set-up. Here cells endogenously expressing and secreting prothrombin, as well as cells induced in disease models, are identified by non-invasive imaging, isolated and cultured. These can be screened ex-vivo through high-throughput luminometry, and additionally in-vivo, through injection into wild-type mice and non-invasive imaging. Findings made in these systems can finally be validated in-vivo using the original D-Insight model. The heat map in the center exemplifies a tissue selective targeting, obtained from a pilot proof-of-concept study (shown in Extended Data Figure).

aberrantly expressed. Additionally, our pilot screening demonstrates that this reporter tool set is suited to uncover tissue-specific regulatory mechanisms that may allow tailored therapeutic targeting. Given the broad expression of prothrombin in various tissues, the reporter model introduced here will likely find wide application to understand

and dissect the role of prothrombin in numerous extravascular pathophysiologicals [36-64].

Biologically, our proof-of-concept observations made here reveal a novel mechanism as to how cancer contributes to a hypercoagulant phenotype. Overexpression of prothrombin perturbs the well-balanced equilibrium between pro- and anticoagulatory activities in the plasma. While the hyperexpression observed in tumors is low level, the cumulative secretion appears to quantitatively exceed the pro-coagulant effects caused by the well-established thrombophilic prothrombin polymorphism F2 20210 G>A [65-66]. Even small tumor lesions can lead to the hypercoagulative state frequently observed in cancer patients, ultimately leading to detrimental consequences including thromboembolic death [53-54]. In view of the evolution of direct oral anticoagulants (DOACs), charting the spatiotemporal dynamics of prothrombin in pathophysiologicals may also reveal hitherto unrecognized advantages of selective therapeutic targeting of the hemostatic system [67-69].

Beyond discovering new functions of the hemostatic system with spatiotemporal resolution, this reporter model is multifunctional. It allows exploring fundamental biological principles of gene regulation and protein targeting, and to identify rheostats involved in the cross-talk between gene expression and availability of a secretory protein. Complementary syngeneic primary cell culture models also enable the deconvolution of regulatory mechanisms in high resolution, and in a scalable high throughput format ex vivo. It thereby represents a valuable resource for uncovering disease-eliciting cues and novel therapeutic vulnerabilities, including their validation by non-invasive imaging in vivo, and vice versa (Figure 5). Ultimately this model is an instructive example for a successful multimodal imaging strategy, which can be simply adapted to numerous other secretory proteins.

Contributions

JN, ST, SK, EK, LKS, LH, LE, DAS performed experiments. JN and ST analyzed and compiled data. HC interpreted data and discussed results. SD conceived and supervised the study, performed experiments and wrote the paper with contributions from all coauthors.

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