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Bayesian Inference Associates Rare *KDR* Variants with Specific Phenotypes in Pulmonary Arterial Hypertension

Running title: *Swietlik et al.; A genotype-phenotype association study in PAH*

Emilia M. Swietlik, MD¹; Daniel Greene, PhD^{2,3}; Na Zhu, PhD^{4,5}; Karyn Megy, PhD^{2,3}; Marcella Cogliano, MSc⁶; Smitha Rajaram, MD⁷; Divya Pandya, MSc¹; Tobias Tilly, MSc¹; Katie A. Lutz, BS⁸; Carrie C.L. Welch, PhD⁴; Michael W. Pauciulo, BS, MBA^{8,9}; Laura Southgate, PhD¹⁰; Jennifer M. Martin, MSt³; Carmen M. Treacy, BSc¹; Christopher J. Penkett, PhD^{2,3}; Jonathan C. Stephens, BSc^{2,3}; Harm J. Bogaard, MD, PhD¹¹; Colin Church, PhD¹²; Gerry Coghlan, MD¹³; Anna W. Coleman, MS⁸; Robin Condliffe, MD¹⁴; Christina A. Eichstaedt, PhD¹⁵⁻¹⁷; Mélanie Eyries, PhD¹⁸; Henning Gall, MD, PhD¹⁹; Stefano Ghio, MD²⁰; Barbara Girerd, PhD²¹; Ekkehard Grünig, MD^{16,17}; Simon Holden, PhD²²; Luke Howard, MD, PhD²³; Marc Humbert, MD, PhD²¹; David G. Kiely, MD¹⁴; Gabor Kovacs, MD^{24,25}; Jim Lordan, PhD²⁶; Rajiv D. Machado, PhD¹⁰; Robert V. MacKenzie Ross, MB, BChir²⁷; Colm McCabe, PhD^{23,28}; Shahin Moledina, MBChB²⁹; David Montani, MD, PhD²¹; Horst Olschewski, MD^{24,25}; Joanna Pepke-Zaba, PhD³⁰; Laura Price, PhD^{23,28}; Christopher J. Rhodes, PhD²³; Werner Seeger, MD¹⁹; Florent Soubrier, MD, PhD¹⁸; Jay Suntharalingam, MD²⁷; Mark R. Toshner, MD^{1,30}; Anton Vonk Noordegraaf, MD¹¹; John Wharton, PhD²³; James M. Wild, PhD⁶; Stephen John Wort, PhD^{23,28}; NIHR BioResource for Translational Research - Rare Diseases³¹; National Cohort Study of Idiopathic and Heritable PAH³²; PAH Biobank Enrolling Centers' Investigators³³; Allan Lawrie, PhD⁶; Martin R. Wilkins, MD²³; Richard C. Trembath, FRCP³⁴; Yufeng Shen, PhD^{5,35}; Wendy K. Chung, MD³⁶; Andrew J. Swift, PhD⁶; William C. Nichols, PhD^{8,9}; Nicholas W. Morrell, MD^{1,3,22,30}; Stefan Gräf, PhD¹⁻³

¹Dept of Medicine, ²Dept of Haematology, Univ of Cambridge; ³NIHR BioResource for Translational Research, Cambridge, UK; ⁴Dept of Pediatrics, ⁵Dept of Systems Biology, ⁶Dept of Biomedical Informatics, Columbia Univ, New York, NY; ⁷Dept of Infection, Immunity & Cardiovascular Disease, Univ of Sheffield; ⁸Sheffield Teaching Hospitals NHS Foundation Trust, Sheffield, UK; ⁹Division of Human Genetics, Cincinnati Children's Hospital Medical Ctr; ¹⁰Dept of Pediatrics, Univ of Cincinnati College of Medicine, Cincinnati, OH; ¹¹Molecular & Clinical Sciences Rsrch Inst, St George's, Univ of London, London, UK; ¹²Dept of Clinical Genetics, Amsterdam UMC, Vrije Universiteit Amsterdam, Amsterdam, the Netherlands; ¹³Golden Jubilee National Hospital, Glasgow; ¹⁴Royal Free Hospital, London; ¹⁵Sheffield Pulmonary Vascular Disease Unit, Royal Hallamshire Hospital, Sheffield, UK; ¹⁶Laboratory for Molecular Genetic Diagnostics, Inst of Human Genetics, Heidelberg Univ; ¹⁷Ctr for Pulmonary Hypertension, Thoraxklinik gGmbH Heidelberg at Heidelberg Univ Hospital; ¹⁸Translational Lung Rsrch Ctr Heidelberg (TLRC), German Ctr for Lung Research (DZL), Heidelberg, Germany; ¹⁹Département de génétique, hôpital Pitié-Salpêtrière, Assistance Publique-Hôpitaux de Paris & UMR_S 1166-ICAN, INSERM, UPMC Sorbonne Universités, Paris, France; ²⁰Univ of Giessen & Marburg Lung Center (UGMLC), member of the German Center for Lung Research (DZL) and of the Excellence Cluster Cardio-Pulmonary Inst (CPI), Giessen, Germany; ²¹Fondazione IRCCS Policlinico San Matteo, Pavia, Italy; ²²Université Paris-Sud, Faculté de Médecine, Université Paris-Saclay; AP-HP, Service de Pneumologie, Centre de référence de l'hypertension pulmonaire; INSERM UMR_S 999, Hôpital Bicêtre, Le Kremlin-Bicêtre, Paris, France; ²³Addenbrooke's Hospital NHS Foundation Trust, Cambridge; ²⁴National Heart & Lung Inst, Imperial College London, London, UK; ²⁵Ludwig Boltzmann Inst for Lung Vascular Rsrch; ²⁶Medical Univ of Graz, Graz, Austria; ²⁷Freeman Hospital, Newcastle upon Tyne; ²⁸Royal United Hospitals Bath NHS Foundation Trust, Bath; ²⁹Royal Brompton & Harefield NHS Foundation Trust; ³⁰Great Ormond Street Hospital, London; ³¹Royal Papworth Hospital NHS Foundation Trust; ³²Univ of Cambridge & Cambridge Univ Hospitals NHS Foundation Trust; ³³www.ipahcohort.com, Cambridge, UK; ³⁴www.pahbiobank.org, Cincinnati, OH; ³⁵Dept of Medical & Molecular Genetics, King's College London, London, UK; ³⁶Columbia Univ Medical Ctr, New York, NY

Correspondence:

Dr Stefan Gräf, PhD
Department of Medicine
University of Cambridge, Level 5
Cambridge University Hospitals, Box 157
Cambridge Biomedical Campus
Cambridge, CB2 0QQ, UK
Tel: (+44) 1223 588036
Email: sg550@cam.ac.uk

Prof. Nicholas W. Morrell, MD
Department of Medicine
University of Cambridge, Level 5
Cambridge University Hospitals, Box 157
Cambridge Biomedical Campus
Cambridge, CB2 0QQ, UK
Tel:(+44) 1223 331666
Email: nwm23@cam.ac.uk

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Abstract

Background - Approximately 25% of patients with pulmonary arterial hypertension (PAH) have been found to harbor rare mutations in disease-causing genes. To identify missing heritability in PAH we integrated deep phenotyping with whole-genome sequencing data using Bayesian statistics.

Methods - We analyzed 13,037 participants enrolled in the NIH BioResource - Rare Diseases (NBR) study, of which 1,148 were recruited to the PAH domain. To test for genetic associations between genes and selected phenotypes of pulmonary hypertension (PH), we used the Bayesian rare-variant association method BeviMed.

Results - Heterozygous, high impact, likely loss-of-function variants in the Kinase Insert Domain Receptor (*KDR*) gene were strongly associated with significantly reduced transfer coefficient for carbon monoxide (KCO, posterior probability (PP)=0.989) and older age at diagnosis (PP=0.912). We also provide evidence for familial segregation of a rare nonsense *KDR* variant with these phenotypes. On computed tomographic imaging of the lungs, a range of parenchymal abnormalities were observed in the five patients harboring these predicted deleterious variants in *KDR*. Four additional PAH cases with rare likely loss-of-function variants in *KDR* were independently identified in the US PAH Biobank cohort with similar phenotypic characteristics.

Conclusions - The Bayesian inference approach allowed us to independently validate *KDR*, which encodes for the Vascular Endothelial Growth Factor Receptor 2 (VEGFR2), as a novel PAH candidate gene. Furthermore, this approach specifically associated high impact likely loss-of-function variants in the genetically constrained gene with distinct phenotypes. These findings provide evidence for *KDR* being a clinically actionable PAH gene and further support the central role of the vascular endothelium in the pathobiology of PAH.

Key words: pulmonary hypertension; genetics, association studies; vascular endothelium; vascular endothelial growth factor receptor; family history; computed tomography

Nonstandard Abbreviations and Acronyms

6MWD	Six minute walking distance	HTN	Systemic hypertension
ACVRL1	Activin-Like Kinase 1	I/HPAH	Idiopathic/Hereditary Pulmonary Arterial Hypertension
APAH	Associated Pulmonary Arterial Hypertension	ICC	Intraclass Correlation Coefficient
APAH:CHD-PAH	PAH associated with congenital heart disease	ILD	Interstitial lung disease
APAH:CTD-PAH	PAH associated with connective tissue disease	KCNK3	Potassium Two Pore Domain Channel Subfamily K Member 3
APAH:HIV-PAH	PAH associated with HIV	KCO	Transfer coefficient for carbon monoxide
APAH:PPH-PAH	PAH associated with portopulmonary hypertension	KDR	Kinase Insert Domain Receptor
AQP1	Aquaporin 1	MAF	Minor Allele Frequency
ASD	Atrial Septal Defect	mPAP	Mean pulmonary artery pressure
ATP13A3	ATPase 13A3	mRAP	Mean right atrial pressure
BA	Bronchial artery	NBR	NIHR BioResource - Rare Diseases
BeviMed	Bayesian Evaluation of Variant Involvement in Mendelian Disease	NIHR	National Institute for Health Research
BF	Bayes factor	NO	Nitric oxide
BHF	British Heart Foundation	NTproBNP	N-terminal pro B-type Natriuretic Peptide
BMI	Body Mass Index	OSA	Obstructive sleep apnoea
BMPR2	Bone Morphogenetic Protein Receptor Type 2	PAD	Peripheral artery disease
BNP	B-type Natriuretic Peptide	PAH	Pulmonary Arterial Hypertension
CAD	Coronary artery disease	PAHBB	US PAH Biobank
CADD	Combined Annotation Dependent Depletion	PAWP	Pulmonary Artery Wedge Pressure
CAVI	Caveolin-1	PH	Pulmonary Hypertension
CHMC	Cincinnati Children's Hospital Medical Center	PH-LD	Pulmonary hypertension associated with lung disease
CI	Cardiac index	PH-LHD	Pulmonary hypertension associated with left heart disease
CKD	Chronic kidney disease	PH-multifactorial	Multifactorial pulmonary hypertension
CO	Cardiac output	pLI	Probability of being loss-of-function intolerant
COPD	Chronic obstructive pulmonary disease	PMAF	Probability that the minor allele count is at least the observed minor allele count
CRP	C reactive protein	PolyPhen-2	Polymorphism Phenotyping v2 score
CT	Computerized Tomography	PP	Posterior probability
CTEPH	Chronic thromboembolic pulmonary hypertension	PVOD/PCH	Pulmonary veno-occlusive disease/ Pulmonary capillary hemangiomatosis
CTPA	Computerised Tomography Pulmonary Angiogram	PVR	Pulmonary vascular resistance
CUMC	Columbia University Medical Center	RDW	Red cell distribution width
CVA	Celebro-vascular accident	RHC	Right heart catheterisation
DM	Diabetes mellitus	Shared	Indicates in which cohort(s) the given variant was identified; i.e. PAH ₍₂₎ , BPD ₍₁₎ means 2 PAH cases and 1 BPD case harbour this variant
eCRF	Electronic Case Report Form	SIFT	Sorting Intolerant From Tolerant prediction score
EIF2AK4	Eukaryotic Translation Initiation Factor 2 Alpha Kinase 4	SMAD ₁	SMAD family member 1
ENG	Endoglin	SMAD ₄	SMAD family member 4
FEV ₁	Forced expiratory volume in one second	SMAD ₉	SMAD family member 9
FHx	Family history	SNV	Single Nucleotide Variants
FPAH	Familial PAH	SOX17	SRY-box 17
FVC	Forced Vital Capacity	SpO ₂ post	Arterial oxygen saturation post exercise
GDF2	Growth Differentiation Factor 2	SpO ₂ pre	Arterial oxygen saturation pre exercise
GerpN	Conservation score of each nucleotide in multi-species alignment	U5416	Sugen
GGO	Ground-glass opacities	SvO ₂	Mixed venous oxygen saturation
gnomAD	Genome Aggregation Database	TBX4	T-Box Transcription Factor 4
GRCh37	Genome Reference Consortium human genome build 37	TLC	Total Lung Capacity
HGVS	Human Genome Variation Society	TOPMed	NHLBI Trans-Omics for Precision Medicine Hb
HGVSc	HGVS notation of coding sequence	TSH	Thyroid Stimulating Hormone
HGVSp	HGVS notation of protein sequence	VEGFR2	Vascular Endothelial Growth Factor Receptor 2
HRCT	High-Resolution Computerised Tomography	WBC	White Blood Cell Count
		WHO FC	World Health Organisation functional class



Introduction

Pulmonary arterial hypertension (PAH) is characterized by pulmonary vascular constriction and obliteration, causing elevation of pulmonary vascular resistance and ultimately, right ventricular failure. Molecular mechanisms such as aberrant angiogenesis¹, metabolic reprogramming and resistance to apoptosis² have been proposed to explain pulmonary vessel remodeling. A

breakthrough in our understanding of the pathobiology underlying PAH was the discovery of heterozygous germline mutations in the gene encoding the bone morphogenetic protein receptor type 2 (*BMPR2*)³, responsible for over 70% of familial PAH (FPAH) cases and 15-20% of idiopathic PAH (IPAH) cases. A smaller proportion (up to 10%) of PAH cases are caused by mutations in activin-like kinase 1 (*ACVRL1*)⁴, endoglin (*ENG*)⁵, SMAD family member 9 (*SMAD9*)⁶, caveolin-1 (*CAVI*), involved in colocalization of BMP receptors⁷, and the potassium channel *KCNK3*, responsible for membrane potential and vascular tone⁸. We recently identified rare pathogenic variants in growth differentiation factor 2 (*GDF2*), which encodes BMP9, a major ligand of the *BMPR2*/*ALK1* receptor complex, as well as rare variants in ATPase 13A3 (*ATP13A3*), aquaporin 1 (*AQP1*) and SRY-box 17 (*SOX17*), and reported a list of additional putative genes potentially contributing to the pathobiology of PAH⁹. Together, the established genes explain approximately 25% of cases with IPAH, allowing their reclassification as heritable PAH (HPAH) cases. To identify additional genes harboring potentially causal rare variants in IPAH cases, we increased the cohort size¹⁰ and deployed a recently developed Bayesian methodology (BeviMed)¹¹ that incorporates phenotypic data to increase the power to detect rare risk variants.

Methods

Figure 1A provides an overview of the analysis strategy. The method details are described in the Supplemental Material. The data of the NBR study have been deposited in the European Genome-Phenome Archive¹⁰. The data from the US PAH Biobank (PAHBB) and the Columbia University Medical Center (CUMC) are available via an application¹².

Patients recruited to the NBR study provided informed consent for genetic analysis and clinical data capture (REC REF: 13/EE/0325); patients recruited by European collaborators

consented to genetic testing and clinical data collection locally. Institutional review boards at Cincinnati Children's Hospital Medical Center (CHMC) and CUMC, and the PAHBB Centers approved the validation cohort studies and written informed consent was obtained at enrollment.

Results

Characterization of study cohorts and tag definition

Whole-genome sequencing was performed in 13,037 participants of the NBR study, of which 1,148 were recruited to the PAH domain¹⁰. The PAH domain included 23 unaffected parents and three cases with an unknown phenotype, which were removed from the analysis (Figure 1B). Of the remaining 1,122 participants, 972 (86.6%) had a clinical diagnosis of IPAH, 73 (6.5%) of HPAH, and 20 (1.8%) were diagnosed with PVOD/PCH. Diagnosis verification revealed that 57 participants (5%) had a diagnosis other than IPAH, HPAH or PVOD/PCH. These cases were subsequently relabelled and moved to the respective tag group for analysis (Table 1). The comprehensive clinical characterisation of the study cohort is shown in Supplemental Table 1. In summary, the median age at diagnosis was 49[35;63] years with a female predominance of 68%. Europeans constituted 84% of the study cohort. Overall survival in the studied population was 97% at one year, 91% at three years and 84% at five years. As expected, there was a significant difference in survival between prevalent and incident cases. In prevalent cases, survival at one, three, and five years was 98%, 93%, and 87%, whereas in incident cases it was 97%, 84%, and 72%, respectively. Median transfer coefficient for carbon monoxide (KCO) in the entire studied population was 71[52;86]% predicted. Cases in the lower tertile or below the KCO threshold of 50% predicted were more commonly male, older at diagnosis, had a current or past history of cigarette smoking and an increased number of cardiorespiratory comorbidities (Supplemental Tables 2, 3, and 4). Survival in these groups was significantly worse than in those with preserved

or mildly reduced KCO (Supplemental Figure 1A-D). After adjusting for confounding factors (age, sex, comorbidities, smoking status and whether the case was prevalent or incident), KCO remained an independent predictor of survival (Supplemental Table 5).

Age at diagnosis was calculated as age at the time of diagnostic right heart catheterisation and was available in all but 10 cases. Patients in the higher age tertile showed more functional impairment despite milder hemodynamics, lower FEV1/FVC ratio and KCO [% predicted], as well as mild emphysematous and fibrotic changes on CT scans (Supplemental Figure 1E and F and Supplemental Table 6).

Rare variants in previously established genes

We identified variants in previously established genes (namely, *BMP2*, *ACVRL1*, *ENG*, *SMAD1*, *SMAD4*, *SMAD9*, *KCNK3*, *TBX4*, *EIF2AK4*, *AQP1*, *ATP13A3*, *GDF2*, *SOX17*) in 271 (24.2%) of the 1,122 cases and interpreted them based on the ACMG standards and guidelines¹³. The majority of these variants have already been described in Gräf *et al.*⁹ (see Supplemental Material).

Rare variant association testing

We used BeviMed to consolidate previously reported PAH genes and to discover novel genotype-phenotype associations. Of note, cases explained by rare deleterious variants in previously established genes were only included for the association testing with the respective disease gene (see Methods). This analysis identified 40 significant gene-tag associations with posterior probability (PP) above 0.75 (Table 2 and Figure 2A). *BMP2*, *TBX4*, *EIF2AK4*, *ACVRL1* and *AQP1* showed the highest association (PP \geq 0.99) but we also confirmed significant associations in the majority of other previously identified genes. Individuals with rare variants in *BMP2*, *TBX4* (high impact), *EIF2AK4* (biallelic) and *SOX17* had a significantly younger age of

disease onset (tag: young age). We also confirmed the association of rare variants in *AQPI* with FPAH (log(BF)=10.023, PP=0.958). The refined phenotype approach corroborated the association between high impact variants in *BMP2* and preserved KCO (KCO higher tertile, log(BF)=99.923, PP=1) together with an association of biallelic *EIF2AK4* mutations with significantly reduced KCO (KCO <50% predicted, log(BF)= 29.741, PP=1).

Under an autosomal dominant mode of inheritance, high impact variants in the Kinase Insert Domain Receptor (*KDR*) were associated with a significantly reduced KCO (KCO lower tertile, log(BF)=11.362, PP=0.989) and older age at diagnosis (tag: old age, log(BF)=9.249, PP=0.912).

Rare high impact variants in the new PAH candidate gene *KDR*



We identified five ultra-rare high impact variants in *KDR* in the study cohort. Ultra-rare variants exist in the general population only at a frequency of less than 1 in 10,000 (0.01%). Four of these were in PAH cases: one frameshift variant in exon 3 of 30 (c.183del, p.Tryp61CysfsTer16), two nonsense variants, one in exon 3 (c.183G>A, p.Trp61Ter) and one in exon 22 (c.3064C>T, p.Arg1022Ter) and one splice acceptor variant in intron 4 of 29 (c.490-1G>A). In addition, one nonsense variant was identified in exon 27 (p.Glu1206Ter) in a non-PAH control subject (Table 3). This latter nonsense variant appears late in the amino acid sequence, in exon 27 of 30, and hence is likely to escape nonsense-mediated decay, but this remains to be studied functionally. All loss-of-function variants were confirmed by Sanger sequencing (Figure 3 and Supplemental Figure 2). Furthermore, 13 PAH cases (1%) and 108 non-PAH controls (0.9%) harbored rare, predicted-deleterious *KDR* missense variants of moderate impact (Figure 3). The missense variant carriers, however, did not exhibit a reduced KCO or older age at diagnosis. Instead, these patients show the opposite trend in KCO (Figure 2B and C). Importantly, seven of the 13 *KDR*

missense variants seen in PAH cases were also detected in several non-PAH controls and thus, are of unknown significance. Furthermore, three of these missense variants co-occurred with a predicted-deleterious variant in an established PAH risk gene (two patients carried also a variant in *BMPR2* and one a variant in *AQP1*).

Clinical characterization of *KDR* mutation carriers

Patients with high impact variants in *KDR* were older and exhibited significantly reduced KCO similar to biallelic *EIF2AK4* mutation carriers and in contrast to *KDR* missense variant and *BMPR2* mutation carriers (Figure 2B and C). Three of the four cases did not have a history of smoking. CT scans for all four patients showed a range of mild lung parenchymal changes (Figure 4). W000229 had evidence of mild mainly subpleural interstitial lung disease (ILD), mild emphysema, and air trapping. W000274 had signs of ILD with traction bronchiectasis in the lower zones, mild air trapping, and mild diffuse ground-glass opacities (GGO) and neovascularity. E001392 showed mild centrilobular GGO in addition to moderate pleural effusion and a trace of air trapping, but no ILD. In these cases, it seemed likely that the observed parenchymal changes contributed to the low KCO. In contrast, E003448 had a low KCO despite only a trace of central nonspecific GGO on the CT images. Comparisons of CT findings between patients harboring deleterious mutations in *BMPR2*, *EIF2AK4*, *KDR*, other PAH risk genes and patients without mutations are presented in Supplemental Table 7. There were no differences in the frequency of comorbidities between patients harboring missense and loss-of-function variants in *KDR* although the frequency of systemic hypertension was high (44%) (Supplemental Table 8). Survival analysis could not be conducted due to the small number of mutation carriers, as well as only two events occurring in this group. Following the death of W000229, his daughter, aged 53, was diagnosed with PAH and had a reduced KCO at 40% predicted. On the CT scan,

mild interstitial fibrosis was observed (Supplemental Figure 3). Sanger sequencing confirmed that father and daughter carried the same deleterious *KDR* nonsense variant p.Trp61Ter (Figure 3B).

Additional *KDR* cases in US PAH cohorts

To seek further evidence for *KDR* as a new candidate gene for PAH, we analyzed subjects recruited to the PAHBB¹² and the CUMC¹⁴ to identify additional patients carrying predicted pathogenic rare variants. Four additional individuals harboring rare high impact *KDR* variants were identified. These comprised, two nonsense variants, one in exon 3 (c.303C>A, p.Tyr101Ter) and one in exon 22 (c.3064C>T, p.Arg1022Ter) and two splice donor variants, one in intron 2 of 29 (c.161+1G>T) and one in intron 5 (c.658+1G>A). Interestingly, the nonsense variant p.Arg1022Ter appeared in both cohorts (Figure 3). Patient-level data for these individuals are summarized in Table 3. Three of the four patients were diagnosed with idiopathic PAH at 72, 65 and 42 years respectively, whereas one patient was diagnosed at age four with PAH associated with double outlet right ventricle. The diffusing capacity of carbon monoxide was available for one patient and was decreased at 35% predicted, with minor pleural scarring in the left upper lobe found on CT imaging. Two out of four patients (50%) harboring a high impact variant in *KDR* had been diagnosed with systemic hypertension.

Discussion

One of the critical steps in identifying novel, causative genes in rare disorders is the discovery of genotype-phenotype associations to inform patient care and outcomes. A pragmatic focus on deeply phenotyped individuals and “smart” experimental design provides additional leverage to identify novel risk variants¹⁵. To deploy this approach in PAH we brought together phenotypic and genetic data using BeviMed¹¹. This Bayesian framework allows the inclusion of prior

information regarding the hypothesis being tested in a flexible manner and compares a range of possible genetic models in a single analysis. To generate case-control labels, we tagged PAH cases with diagnostic labels and stratified them by age at diagnosis and KCO. Analyses were then performed to identify associations between tags and ultra-rare gene variants under dominant and recessive modes of inheritance and different variant impact categories.

Our BeviMed analysis provided strong statistical evidence of an association between ultra-rare, high impact variants in *KDR* and PAH with significantly reduced KCO and older age at diagnosis under a dominant mode of inheritance. Strikingly, likely loss-of-function variants in *KDR* exist in the general population with a frequency of only 4-7 per 100,000 (see Table 4). In contrast, we identified four PAH cases in the NBR cohort which equates to almost 2 in 1,000. Additionally, the statistical constraint metrics provided by gnomAD¹⁶ strongly suggest that loss-of-function variants in *KDR* are not tolerated (pLI = 1; o/e = 0.15 (0.09 - 0.25)). Besides the statistical evidence, we also identified one additional case with a family history, which together with a recently published case report of two families in which loss-of-function variants in *KDR* segregated with PAH and significantly reduced KCO¹⁷, amounts to three reported familial cases with a distinct phenotype.

VEGFR2, which is encoded by *KDR*¹⁸, binds VEGFA, a critical growth factor for physiological and pathological angiogenesis in vascular endothelial cells. In mice, even though VegfA haploinsufficiency is embryonically lethal¹⁹, heterozygosity of its receptor, Vegfr2, is compatible with life and unperturbed vascular development²⁰. The role of VEGF signaling in the pathogenesis of PAH has been an area of intense interest since increased expression of VEGF, VEGFR1 and VEGFR2 were reported in rat lung tissue in response to acute and chronic hypoxia²¹. An increase in lung VEGF has also been reported in rats with PH following

monocrotaline exposure²². In humans, VEGF-A is highly expressed in plexiform lesions in patients with IPAH²³. In addition, inhibition of VEGF signaling by SU5416 (sugen) combined with chronic hypoxia triggers severe angioproliferative PH²⁴. SU5416, a small-molecule inhibitor of the tyrosine kinase segment of VEGF receptors, inhibits VEGFR1²⁵ and VEGFR2²⁶ causing endothelial cell apoptosis, loss of lung capillaries and emphysema²⁷. Further evidence supporting the role of VEGF inhibition in the pathobiology of PAH comes from reports of PH in patients treated with bevacizumab²⁸ and the multi-tyrosine kinase inhibitors^{29,30}. Mutations in *KDR* have also been linked to congenital heart diseases. Bleyl *et al.* reported that *KDR* might be a candidate for familial total anomalous pulmonary venous return³¹. In addition, haploinsufficiency at the *KDR* locus has also been associated with tetralogy of Fallot³². We identified one patient in the CUMC cohort with PAH associated with congenital heart disease harboring a *KDR* likely protein-truncating splice donor variant (c.161+1G>T).

In the present study, we highlight that deep clinical phenotyping, in combination with genotype data, can improve the identification of novel disease risk genes and disease subtypes. *KDR* was already identified as a possible candidate gene, which did not achieve genome-wide significance, in our previous rare variant association study⁹. In combination with deep phenotyping data, *KDR* reached in the present study a significance level comparable to the most commonly affected genes in PAH. Reduced KCO, which reflects impairment of alveolar-capillary membrane function, has been noted in the analysis of early PAH registry data³³ to be an independent predictor of survival. Decreased KCO was also found in patients with PVOD/PCH with or without biallelic *EIF2AK4* mutations³⁴. Although some reduction in KCO is one of the typical features of pulmonary vascular disease, PVOD patients show the lowest KCO values when compared to IPAH or CTEPH. In contrast, KCO is relatively preserved in *BMPR2*

mutation carriers³⁵. Strong association with survival and a link with other causative mutations makes the KCO phenotype particularly attractive for stratification in genetic studies.

As lung disease should always be taken under consideration as a cause of low KCO, we applied the World Symposium on PH criteria³⁶ to exclude lung disease as a cause of PH: TLC $\geq 70\%$ pred., FVC $\geq 70\%$ pred., FEV1 $\geq 60\%$ pred., and no severe fibrosis and/or emphysema on chest CT. None of the cases carrying a high impact variant in *KDR* met these criteria, although two of the four patients did show evidence of early ILD. Another potential reason for low KCO in the PAH population is the diagnosis of PVOD/PCH³⁷. Careful analysis of CT scans and clinical data did not reveal convincing evidence for this diagnosis in *KDR* mutation carriers.

Cigarette smoking is a well-known factor leading to the decrease of KCO. Only one of the four *KDR* high impact variant carriers had a significant 15 pack-years smoking history, but with no signs of emphysema on CT. These findings suggest that loss-of-function variants in *KDR* are associated with a form of PAH characterized by a range of lung parenchymal abnormalities, including small airways disease, emphysema and ILD, as two of the four patients harboring a high impact variant in *KDR* had mild fibrotic lung changes. Notably, patients with mutations in other PAH risk genes, or those without the identified genetic mutation, showed less than 10% incidence of fibrotic changes on CT imaging. Further larger studies are needed to determine the full range of lung parenchymal abnormalities in PAH cases with deleterious variants in *KDR*.

In this study, we have assumed that PAH is a monogenic condition, which is caused by either deleterious heterozygous or biallelic variants in a single gene. This assumption, although widely supported by the literature, may not be entirely accurate. Alternatively, some cases of PAH might represent an oligogenic inheritance involving two or more genes. Although not statistically explored in the current analysis we found a total of 22 PAH cases carrying

deleterious variants in more than one PAH gene. These variants could contribute as genetic modifiers, impacting penetrance and/or expressivity. In this analysis, we have explored only a limited number of clinical phenotypes. Further studies with larger numbers of phenotypic tags derived from clinical and molecular data will increase the power to detect new associations. Finally, KCO measurements were missing for a proportion of patients which could introduce a selection bias, although all the deleterious variants in *KDR* had phenotypic data available in the UK cohort.

In summary, this study shows that deep phenotyping enables patient stratification into subgroups with shared pathobiology and with increased power to detect new genotype-phenotype associations. We provide statistical evidence for an association between high impact, likely loss-of-function variants in *KDR* and significantly decreased KCO and later disease onset, further supported by familial segregation.

Appendix:

NIHR BioResource for Translational Research - Rare Diseases / National Cohort Study of Idiopathic and Heritable PAH collaborator: Stephen Abbs, Lara Abulhoul, Julian Adlard, Munaza Ahmed, Timothy J Aitman, Hana Alachkar, David J Allsup, Philip Ancliff, Richard Antrobus, Ruth Armstrong, Gavin Arno, Sofie Ashford, William J Astle, Anthony Attwood, Paul Aurora, Christian Babbs, Chiara Bacchelli, Tamam Bakchoul, Siddharth Banka, Tadbir Bariana, Julian Barwell, Joana Batista, Helen E Baxendale, Phil L Beales, David L Bennett, Agnieszka Bierzynska, Tina Biss, Maria A K Bitner-Glindzicz, Graeme C Black, Marta Bleda, Iulia Blesneac, Detlef Bockenhauer, Harm Bogaard, Sara Boyce, John R Bradley, Gerome Breen, Paul Brennan, Carole Brewer, Matthew Brown, Andrew C Browning, Michael J Browning, Rachel J Buchan, Matthew S Buckland, Teofila Bueser, Carmen Bugarin Diz, John Burn, Siobhan O Burns, Oliver S Burren, Nigel Burrows, Carolyn Campbell, Gerald Carr-White, Keren Carss, Ruth Casey, Mark J Caulfield, Jenny Chambers, John Chambers, Melanie M Y Chan, Floria Cheng, Patrick F Chinnery, Manali Chitre, Martin T Christian, Colin Church, Jill Clayton-Smith, Maureen Cleary, Naomi Clements Brod, Gerry Coghlan, Elizabeth Colby, Trevor R P Cole, Janine Collins, Peter W Collins, Cecilia J Compton, Robin Condliffe, H Terence Cook, Stuart Cook, Nichola Cooper, Paul A Corris, Nicola S Curry, Matthew J Daniels, Mehul Dattani, Louise C Daugherty, John Davis, Anthony De Soyza, Sri V V Deevi, Timothy Dent, Charu Deshpande, Eleanor F Dewhurst, Peter H Dixon, Sofia Douzgou, Kate Downes, Anna M Drazyk, Elizabeth Drewe, Daniel Duarte, Tina Dutt, J David M Edgar, Karen Edwards, William Egner, Melanie N Ekani, Perry Elliott, Wendy N Erber, Marie Erwood, Maria C Estiu, Dafydd Gareth

Evans, Gillian Evans, Tamara Everington, Mèlanie Eyries, Hiva Fassihi, Remi Favier, Debra Fletcher, Frances A Flinter, R Andres Floto, Tom Fowler, James Fox, Amy J Frary, Courtney E French, Kathleen Freson, Mattia Frontini, Abigail Furnell, Daniel P Gale, Henning Gall, Vijeya Ganesan, Michael Gattens, Stefano Ghio, Hossein-Ardeschir Ghofrani, J Simon R Gibbs, Kate Gibson, Kimberly C Gilmour, Barbara Girerd, Nicholas S Gleadall, Sarah Goddard, Keith Gomez, Pavels Gordins, David Gosal, Stefan Gräf, Jodie Graham, Luigi Grassi, Daniel Greene, Lynn Greenhalgh, Andreas Greinacher, Paolo Gresele, Philip Griffiths, Sofia Grigoriadou, Detelina Grozeva, Mark Gurnell, Scott Hackett, Charaka Hadinnapola, Rosie Hague, William M Hague, Matthias Haimel, Matthew Hall, Helen L Hanson, Eshika Haque, Kirsty Harkness, Andrew R Harper, Claire L Harris, Daniel Hart, Ahamad Hassan, Grant Hayman, Alex Henderson, Archana Herwadkar, Jonathan Hoffman, Simon Holden, Rita Horvath, Henry Houlden, Arjan C Houweling, Luke S Howard, Fengyuan Hu, Gavin Hudson, Aarnoud P Huissoon, Marc Humbert, Matthew Hurles, Melita Irving, Louise Izatt, Roger James, Sally A Johnson, Stephen Jolles, Jennifer Jolley, Dragana Josifova, Neringa Jurkute, Mary A Kasanicki, Hanadi Kazkaz, Rashid Kazmi, Peter Kelleher, Anne M Kelly, Wilf Kelsall, Carly Kempster, David G Kiely, Nathalie Kingston, Nils Koelling, Myrto Kostadima, Gabor Kovacs, Ania Koziell, Roman Kreuzhuber, Taco W Kuijpers, Ajith Kumar, Dinakantha Kumararatne, Manju A Kurian, Michael A Laffan, Fiona Laloo, Michele Lambert, Hana Lango Allen, Allan Lawrie, D Mark Layton, Claire Lentaigne, Tracy Lester, Adam P Levine, Rachel Linger, Hilary Longhurst, Lorena E Lorenzo, Eleni Louka, Paul A Lyons, Rajiv D Machado, Robert V MacKenzie Ross, Bella Madan, Eamonn R Maher, Jesmeen Maimaris, Samantha Malka, Sarah Mangles, Rutendo Mapeta, Kevin J Marchbank, Stephen Marks, Hugh S Markus, Hanns-Ulrich Marschall, Andrew Marshall, Jennifer Martin, Mary Mathias, Emma Matthews, Heather Maxwell, Paul McAlinden, Mark I McCarthy, Harriet McKinney, Stuart Meacham, Adam J Mead, Karyn Megy, Sarju G Mehta, Michel Michaelides, Carolyn Millar, Shehla N Mohammed, Shahin Moledina, David Montani, Anthony T Moore, Nicholas W Morrell, Monika Mozere, Keith W Muir, Andrew D Mumford, Andrea H Nemeth, William G Newman, Michael Newnham, Sadia Noorani, Paquita Nurden, Jennifer O'Sullivan, Samya Obaji, Chris Odhams, Steven Okoli, Andrea Olschewski, Horst Olschewski, Kai Ren Ong, S Helen Oram, Elizabeth Ormondroyd, Willem H Ouwehand, Claire Palles, Sofia Papadia, Soo-Mi Park, David Parry, Smita Patel, Joan Paterson, Andrew Peacock, Simon H Pearce, Kathelijne Peerlinck, Christopher J Penkett, Joanna Pepke-Zaba, Romina Petersen, Clarissa Pilkington, Kenneth E S Poole, Bethan Psaila, Angela Pyle, Richard Quinton, Shamima Rahman, Anupama Rao, F Lucy Raymond, Paula J Rayner-Matthews, Augusto Rendon, Tara Renton, Christopher J Rhodes, Andrew S C Rice, Alex Richter, Leema Robert, Irene Roberts, Sarah J Rose, Robert Ross-Russell, Catherine Roughley, Noemi B A Roy, Deborah M Ruddy, Omid Sadeghi-Alavijeh, Moin A Saleem, Nilesh Samani, Crina Samarghitean, Alba Sanchis-Juan, Ravishankar B Sargur, Robert N Sarkany, Simon Satchell, Sinisa Savic, Genevieve Sayer, John A Sayer, Laura Scelsi, Andrew M Schaefer, Sol Schulman, Richard Scott, Marie Scully, Claire Searle, Werner Seeger, Arjune Sen, W A Carrock Sewell, Denis Seyres, Neil Shah, Olga Shamardina, Susan E Shapiro, Adam C Shaw, Keith Sibson, Lucy Side, Ilenia Simeoni, Michael A Simpson, Matthew C Sims, Suthesh Sivapalaratnam, Damian Smedley, Katherine R Smith, Kenneth G C Smith, Katie Snape, Nicole Soranzo, Florent Soubrier, Laura Southgate, Olivera Spasic-Boskovic, Simon Staines, Emily Staples, Hannah Stark, Jonathan Stephens, Kathleen E Stirrups, Alex Stuckey, Jay Suntharalingam, Emilia M Swietlik, Petros Syrris, R Campbell Tait, Kate Talks, Rhea Y Y Tan, Jenny C Taylor, John M Taylor, James E Thaventhiran, Andreas C Themistocleous, David Thomas, Ellen Thomas, Moira

J Thomas, Patrick Thomas, Kate Thomson, Adrian J Thrasher, Chantal Thys, Tobias Tilly, Marc Tischkowitz, Catherine Titterton, Cheng-Hock Toh, Ian P Tomlinson, Mark Toshner, Matthew Traylor, Carmen Treacy, Paul Treadaway, Richard Trembath, Salih Tuna, Ernest Turro, Philip Twiss, Tom Vale, Chris Van Geet, Natalie van Zuydam, Anthony M Vandersteen, Marta Vazquez-Lopez, Julie von Ziegenweidt, Anton Vonk Noordegraaf, Annette Wagner, Quinten Waisfisz, Neil Walker, Suellen M Walker, James S Ware, Hugh Watkins, Christopher Watt, Andrew R Webster, Lucy Wedderburn, Wei Wei, Steven B Welch, Julie Wessels, Sarah K Westbury, John-Paul Westwood, John Wharton, Deborah Whitehorn, James Whitworth, Andrew O M Wilkie, Martin R Wilkins, Catherine Williamson, Brian T Wilson, Edwin K S Wong, Nicholas Wood, Yvette Wood, Christopher Geoffrey Woods, Emma R Woodward, Stephen J Wort, Austen Worth, Michael Wright, Katherine Yates, Patrick F K Yong, Timothy Young, Ping Yu, Patrick Yu-Wai-Man, Eliska Zlamalova.

PAH Biobank Enrolling Centers' Investigators: Russel Hirsch, MD; R. James White, MD, PhD; Marc Simon, MD; David Badesch, MD; Erika Rosenzweig, MD; Charles Burger, MD; Murali Chakinala, MD; Thenappan Thenappan, MD; Greg Elliott, MD; Robert Simms, MD; Harrison Farber, MD; Robert Frantz, MD; Jean Elwing, MD; Nicholas Hill, MD; Dunbar Ivy, MD; James Klinger, MD; Steven Nathan, MD; Ronald Oudiz, MD; Ivan Robbins, MD; Robert Schilz, DO, PhD; Terry Fortin, MD; Jeffrey Wilt, MD; Delphine Yung, MD; Eric Austin, MD; Ferhaan Ahmad, MD, PhD; Nitin Bhatt, MD; Tim Lahm, MD; Adaani Frost, MD; Zeenat Safdar, MD; Zia Rehman, MD; Robert Walter, MD; Fernando Torres, MD; Sahil Bakshi, DO; Stephen Archer, MD; Rahul Argula, MD; Christopher Barnett MD; Raymond Benza MD; Ankit Desai MD; Veeranna Maddipati MD.

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Table 1. Definitions of labels and the number of unrelated cases and controls in the rare variant association analysis with BeviMed. See paragraph on "Number of PAH domain samples in the analysis" in the Supplemental Material for more details.

Tag	Tag description	Cases	Controls	Excluded relatives
PH	Individuals with mPAP > 25 mmHg	1112	9134	2786
PAH	Patients with one of the following diagnoses: IPAH, HPAH, PVOD, PCH, APAH:CHD-PAH, APAH:CTD-PAH, APAH:HIV-PAH, APAH:PH-PAH	1085	9134	2786
I/HPAH	Patients with a clinical diagnosis of IPAH or HPAH	1036	9134	2786
IPAH	Patients with a clinical diagnosis of IPAH	972	9134	2785
HPAH	Patients with a clinical diagnosis of HPAH	67	9136	2779
PVOD/PCH	Patients with a clinical diagnosis of PVOD/PCH	20	9136	2778
I/HPAH/PVOD/PCH	Patients with one of the following diagnoses: IPAH, HPAH, PVOD, PCH	1056	9134	2786
FPAH	Patients with one of the following diagnoses: IPAH, HPAH, PVOD, PCH and a positive family history	80	9136	2781
APAH	Patients with one of the following diagnoses: APAH:CHD-PAH, APAH:CTD-PAH, APAH:HIV-PAH, APAH:PH-PAH	29	9136	2778
APAH: CHD-PAH	Patients with PAH associated with congenital heart disease	17	9136	2778
APAH: CTD-PAH	Patients with PAH associated with connective tissue disease	10	9136	2778
APAH: PoPH	Patients with PAH associated with portopulmonary hypertension	1	9136	2778
APAH: HIV-PAH	Patients with PAH associated with HIV	1	9136	2778
PH-LHD	Patients with pulmonary hypertension associated with left heart disease (Group 2)	7	9136	2778
PH-LD	Patients with pulmonary hypertension associated with lung disease (Group 3)	8	9136	2778
CTEPH	Chronic thromboembolic pulmonary hypertension (Group 4)	6	9136	2778
PH-multifactorial	Multifactorial pulmonary hypertension (Group 5)	6	9136	2778
young age	Lower age tertile (<40.8 years)	378	9136	2785
middle age	Middle age tertile (40.8 - 58.6 years)	376	9134	2779
old age	Higher age tertile (>58.6 years)	355	9136	2778
low KCO	KCO <50% pred.	152	9136	2778
KCO lower tertile	KCO <60% pred.	211	9136	2778
KCO middle tertile	KCO 60-80% pred.	215	9136	2778
KCO higher tertile	KCO >80% pred.	215	9134	2779

Abbreviations: mPAP - mean pulmonary artery pressure, PH - pulmonary hypertension, PAH - pulmonary arterial hypertension, I/HPAH - Idiopathic/Hereditary Pulmonary Arterial Hypertension, PVOD - Pulmonary veno-occlusive disease, PCH - Pulmonary capillary haemangiomas, APAH - Associated Pulmonary Arterial Hypertension, CHD - Congenital Heart Disease, CTD - Connective Heart Disease, LHD - Left Heart Disease, LD - Lung Disease, CTEPH - Chronic Thromboembolic Pulmonary Hypertension, KCO - transfer coefficient for carbon monoxide.

Table 2. BeviMed analysis results. Posterior probabilities and Bayes Factors (BF) of gene-tag associations (prior probability $\pi = 0.001$). The BF is the ratio between the probabilities of the data under H1 and under H0. The observed data are BF times more likely under H1 than under H0, and so the larger the BF, the stronger the support in the data for H1 compared with H0. The "High" category comprises only variants of high impact, including loss-of-function variants and large deletions; the "Moderate" category contains variants of moderate impact, including missense variants or variants of consequence type "non_coding_transcript_exon_variant"; the combined category "Moderate and High" includes both respective consequence types.

Gene	Transcript	Tag	log(Bayes Factor)	Posterior probability	Consequence type	Mode of inheritance
<i>BMP2</i>	ENST00000374580	I/HPAH	265.762	1.000	High	dominant
<i>BMP2</i>	ENST00000374580	PAH	265.639	1.000	High	dominant
<i>BMP2</i>	ENST00000374580	I/HPAH/PVOD/PCH	263.481	1.000	High	dominant
<i>BMP2</i>	ENST00000374580	PH	262.625	1.000	High	dominant
<i>BMP2</i>	ENST00000374580	young age	149.576	1.000	Moderate and high	dominant
<i>BMP2</i>	ENST00000374580	HPAH	149.091	1.000	Moderate and high	dominant
<i>BMP2</i>	ENST00000374580	FPAH	147.822	1.000	Moderate and high	dominant
<i>BMP2</i>	ENST00000374580	IPAH	144.582	1.000	High	dominant
<i>BMP2</i>	ENST00000374580	KCO higher tertile	99.923	1.000	High	dominant
<i>BMP2</i>	ENST00000374580	middle age	63.119	1.000	Moderate and high	dominant
<i>BMP2</i>	ENST00000374580	KCO middle tertile	52.706	1.000	Moderate and high	dominant
<i>EIF2AK4</i>	ENST00000263791	low KCO	29.741	1.000	Moderate and high	recessive
<i>EIF2AK4</i>	ENST00000263791	KCO lower tertile	26.247	1.000	Moderate and high	recessive
<i>TBX4</i>	ENST00000240335	I/HPAH	23.783	1.000	High	dominant
<i>TBX4</i>	ENST00000240335	I/HPAH/PVOD/PCH	23.549	1.000	High	dominant
<i>TBX4</i>	ENST00000240335	PAH	23.141	1.000	High	dominant
<i>TBX4</i>	ENST00000240335	PH	22.877	1.000	High	dominant
<i>EIF2AK4</i>	ENST00000263791	young age	20.547	1.000	Moderate and high	recessive
<i>TBX4</i>	ENST00000240335	IPAH	19.990	1.000	High	dominant
<i>EIF2AK4</i>	ENST00000263791	I/HPAH/PVOD/PCH	15.718	1.000	Moderate and high	recessive
<i>ACVRL1</i>	ENST00000388922	HPAH	15.501	1.000	Moderate and high	dominant
<i>EIF2AK4</i>	ENST00000263791	PAH	15.407	1.000	Moderate and high	recessive
<i>EIF2AK4</i>	ENST00000263791	PH	15.071	1.000	Moderate and high	recessive

EIF2AK4	ENST00000263791	PVOD/PCH	14.441	0.999	Moderate and high	recessive
AQPI	ENST00000311813	HPAH	12.075	0.994	Moderate	dominant
EIF2AK4	ENST00000263791	FPAH	11.858	0.993	High	recessive
TBX4	ENST00000240335	young age	11.500	0.990	High	dominant
AQPI	ENST00000311813	I/HPAH	11.466	0.990	Moderate and high	dominant
KDR	ENST00000263923	KCO lower tertile	11.362	0.989	High	dominant
AQPI	ENST00000311813	I/HPAH/PVOD/PCH	11.291	0.988	Moderate and high	dominant
AQPI	ENST00000311813	PAH	11.047	0.984	Moderate and high	dominant
AQPI	ENST00000311813	PH	10.791	0.980	Moderate and high	dominant
AQPI	ENST00000311813	FPAH	10.023	0.958	Moderate	dominant
KDR	ENST00000263923	old age	9.249	0.912	High	dominant
GDF2	ENST00000249598	I/HPAH	9.091	0.899	Moderate and high	dominant
BMPR2	ENST00000374580	old age	8.913	0.881	High	dominant
GDF2	ENST00000249598	I/HPAH/PVOD/PCH	8.775	0.866	Moderate and high	dominant
SOX17	ENST00000297316	young age	8.554	0.839	Moderate and high	dominant
GDF2	ENST00000249598	PAH	8.478	0.828	Moderate and high	dominant
ATP13A3	ENST00000439040	KCO higher tertile	8.035	0.755	High	dominant

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Table 3. Gene changes for IPAH patients harboring likely loss-of-function variants in the KDR gene. None of the KDR variants have previously been reported in gnomAD, ExAC or internal controls. HGVS_c notations are based on transcript sequence ENST00000263923.4. HGVS_p notations are based on the amino acid sequence ENSP00000263923.4.

Cohort	UK					US			
	W000229	W000229.d	E003448	W000274	E001392	CUMC-JM161	CCHMC-12-190	CCHMC-19-023	CCHMC-27-015
Exon	3/30	3/30		22/30	3/30	2/30	3/30	5/30	22/30
HGVS _c	c.183G>A	c.183G>A	c.490-1G>A	c.3064C>T	c.183del	c.161+1G>T	c.303C>A	c.658+1G>A	c.3064C>T
HGVS _p	p.Trp61Ter	p.Trp61Ter	-	p.Arg1022Ter	p.Trp61CysfsTer16		p.Tyr101Ter		p.Arg1022Ter
Consequence type	stop gained	stop gained	splice acceptor variant	stop gained	frameshift variant	splice donor variant	stop gained	stop gained	stop gained
Shared	PAH(1)	PAH(1)	PAH(1)	PAH(1)	PAH(1)	PAH(1)	PAH(1)	PAH(1)	PAH(1)
gnomAD	NA	NA	NA	NA	NA	NA	NA	NA	NA
CADD PHRED v1.3	38	38	26	37	35	26.4	38	24.3	37
SIFT	-	-	-	-	-	-	-	-	-
PolyPhen	-	-	-	-	-	-	-	-	-
GerpN	5.93	5.93	5.75	5.95	5.93	5.83	5.48	5.8	5.95
Ancestry	European	European	European	European	European	East-Asian	European	European	European
Sex	male	female	female	male	female	female	male	female	female
Diagnosis	IPAH	IPAH	IPAH	IPAH	IPAH	APAH-CHD secondary to double outlet RV	IPAH	IPAH	IPAH
Age at diagnosis [years]	71	53	62	67	61	4	72	65	42
WHO FC	2	3	3	3	3	2	NA	NA	NA

6MWD [m]	472	202	422	660	180	NA	380	NA	245
SpO₂ pre [%]	95	96	97	98	97	NA	NA	NA	NA
SpO₂ post [%]	86	87	86	NA	91	NA	NA	NA	NA
FEV₁ [% pred.]	116	70	90	83	67.3	85%	NA	77%	NA
FVC [% pred.]	115	76	94	91	72.8	92%	NA	83%	NA
TLC [% pred.]	NA	69	NA	NA	NA	NA	NA	65%	NA
KCO [% pred.]	44	40	46	46	55.2	NA	NA	35%*	NA
Smoking history	Never	Never	Never	Ex-smoker	Never	Never	Never	Ex-smoker	Never
mRAP [mmHg]	5	13	8	8	3	NA	5	29	14
mPAP [mmHg]	62	45	57	41	44	NA	49	66	60
PAWP [mmHg]	4	5	15	12	9	NA	5	16	15
CO [L/min]	3.6	3.3	4.58	5.97	5.23	NA	4.33	1.8	4.6
PVR	16.11	12.12	9.17	4.86	6.69	NA	NA	27.9	9.8
Comorbidities	hyperlipidemia, HTN, DM type 2	DM type 2, OSA, pulmonary fibrosis	HTN, hypothyroidism	DM type 2	CAD, DM type 2	No	HTN, hyperlipidemia,	HTN, hypothyroidism, OA	Obesity, CAD, DM type 2, hypothyroidism
Family history	Yes, daughter	Yes, father	No	No	No	No	No	No	No
Status	dead	alive	alive	alive	dead	alive	alive	alive	alive

Abbreviations: KDR - Kinase insert domain receptor, WHO FC - World Health Organization functional class, 6MWD - 6-minute walk distance, SpO₂ - arterial oxygen saturation, mRAP - mean right atrial pressure, mPAP - mean pulmonary artery pressure, mPAWP - mean pulmonary artery wedge pressure, CO - cardiac output, PVR - pulmonary vascular resistance, FEV₁ - forced expiratory volume in 1 sec, FVC - forced vital capacity, KCO - transfer factor coefficient for carbon monoxide, HTN - systemic hypertension, CAD - coronary artery disease, OA - osteoarthritis, DM - diabetes mellitus, OSA - obstructive sleep apnea, ASD - atrial septal defect, VSD - ventricular septal defect. * DLCO % predicted.

Table 4. Comparison of high impact likely loss-of-function variants in *KDR* in the Human large-scale sequencing reference populations gnomAD and TOPMed with the NBR non-PAH controls and PAH cases.

Large-scale sequencing population	High impact LoF variants in <i>KDR</i>	Individuals	Alleles	Frequency
gnomAD (v2.1)*	20	141,456	282,912	0.000071
gnomAD (v3)	10	71,702	143,404	0.000070
TOPMed	5	62,784	125,568	0.000040
NBR non-PAH controls	1	11,889	23,778	0.000042
NBR PAH cases	4	1,122	2,244	0.001783

**KDR* constraint metrics: pLI = 1; o/e = 0.15 (0.09 - 0.25); exp(LoF) = 73; obs(LoF) = 11

Abbreviations: gnomAD - The Genome Aggregation Database, TOPMed - The Trans-Omics for Precision Medicine program, *KDR* - kinase insert domain receptor, NBR - NIH BioResource - Rare Diseases, LoF - loss of function



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Figure Legends:

Figure 1. Design of the genetic association study. A, Overview of the analytical approach. Using deep phenotyping, data tags were assigned to patients who shared phenotypic features. Rare sequence variants, called from whole-genome sequencing data, were filtered and explained cases were labeled. BeviMed was applied to a set of unrelated individuals to estimate the posterior probability of gene-tag associations. B, Consort diagram summarizing the size of the study cohort. C, Schematic representation of the definition of cases, exemplified by the KCO lower tertile tag. Cases were defined as individuals carrying a particular tag, whereas patients with

missing information or those without a tag were removed from the gene-tag association testing. Individuals from non-PAH domains served as controls. KCO - transfer coefficient for carbon monoxide, MAF - minor allele frequency.

Figure 2. Rare variant association study results revealing established and novel genotype-phenotype links. A, Figure showing phenotype tags on the x-axis and corresponding posterior probability of genotype-phenotype association on the y-axis, as calculated by BeviMed. The definitions of the tags are listed in Table 1. Shape and colour of points indicate the mode of inheritance and impact/consequence type of variants driving the association. Box-and-whisker plots showing the distribution of (B) the transfer coefficient for carbon monoxide (KCO) and (C) the age at diagnosis stratified by genotype across the PAH domain. The two-tailed Wilcoxon signed-rank test was used to determine differences in the medians of the distributions, which are indicated by the bars at the top of the figures providing the respective p-values. Abbreviations: bial. - biallelic, lof - loss-of-function, mis. - missense.

Figure 3. Summary of rare single nucleotide variants (SNVs) and small insertions and deletions (indels) identified in the novel PAH candidate gene KDR. A, Only rare predicted deleterious variants in KDR are shown (MAF<1/10,000 and CADD≥10). SNVs and indels are represented by colored lollipops on top of the protein sequence. The domain annotations were retrieved from Uniprot (accession number P35968). Lollipop colors indicate the consequence type and sizes represent the variant frequency within a cohort. Missense variants that are predicted to be deleterious (SIFT) and damaging (PolyPhen-2) are colored in red, otherwise in yellow (i.e. SIFT and PolyPhen-2 disagree). High impact variants are labelled with the respective HGVS notation.

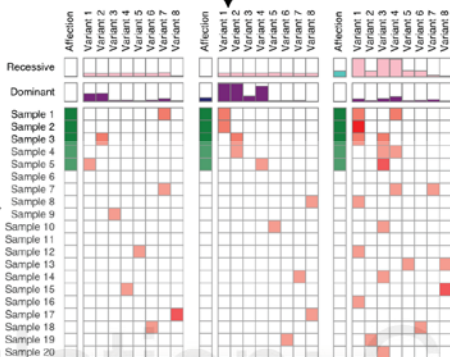
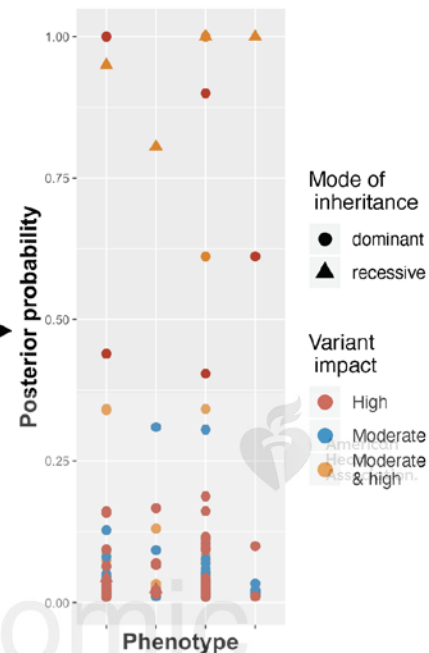
The number of variants by predicted consequence type and cohort is provided in the table. B, Familial segregation of KDR nonsense variant c.183G>A (p.Trp61*) with PAH (i.e. reduced KCO and late onset) from father (W000229) to daughter (W000229.d). Sanger sequencing results are shown in the chromatograms.

Figure 4. Chest computerized tomography (CT) scans of patients carrying high impact KDR mutations. A, Axial image of CT pulmonary angiogram at the level of the right ventricle (RV) moderator band, showing flattening of interventricular septum, leftwards bowing of the interatrial septum and the enlargement of the right atrium (RA) and RV, indicative of RV strain; bilateral pleural effusion, larger on the right side. B, Axial image of a pulmonary CT angiogram demonstrating enlarged pulmonary artery and mild central lung ground-glass opacity (GGO). C, Axial high-resolution CT slice of the chest in the lung window showing a trace of non-specific GGO with a central distribution. D, Coronal image showing the trace of central GGO and enlarged central pulmonary arteries. Axial high-resolution CT slice of the chest in the lung window showing apical subpleural fibrosis (E), and very minor subpleural fibrosis at the lung bases (F). Axial high-resolution CT slice of the chest in the lung window showing subpleural GGO at apical level (G), and mild GGO at mid-thoracic level (H). Patients: E001392 (A, B), E003448 (C, D), W000229 (E, F), W000274 (G,H).

A**Phenotypes****Definition of case-control labels ('tags')****Genotypes**

protein-coding
MAF < 1/10,000
CADD >= 10

label explained cases

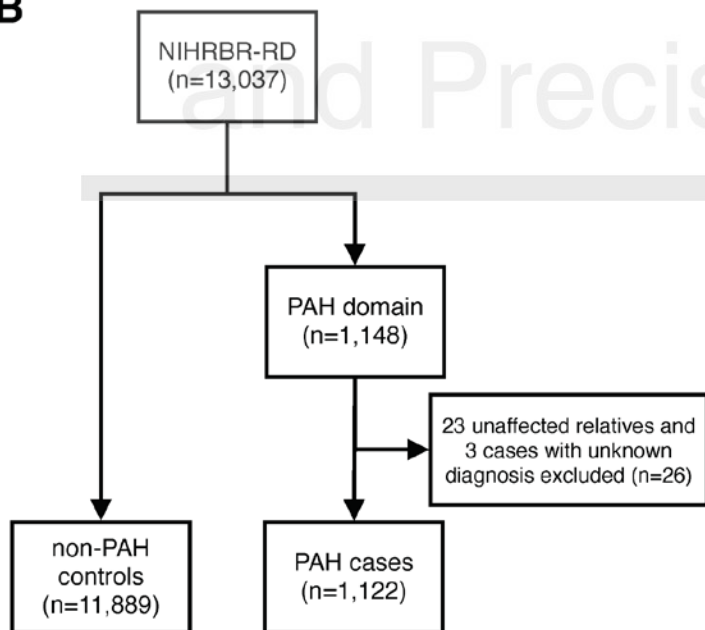
**BeviMed****C**

Mode of inheritance

- dominant
- ▲ recessive

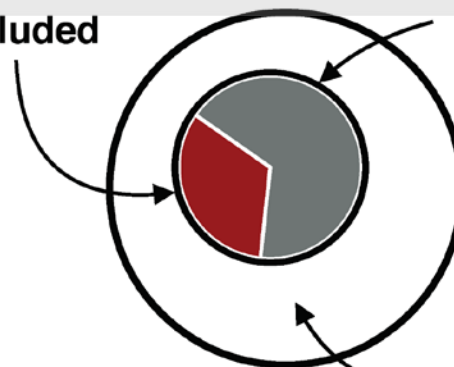
Variant impact

- High
- Moderate
- Moderate & high

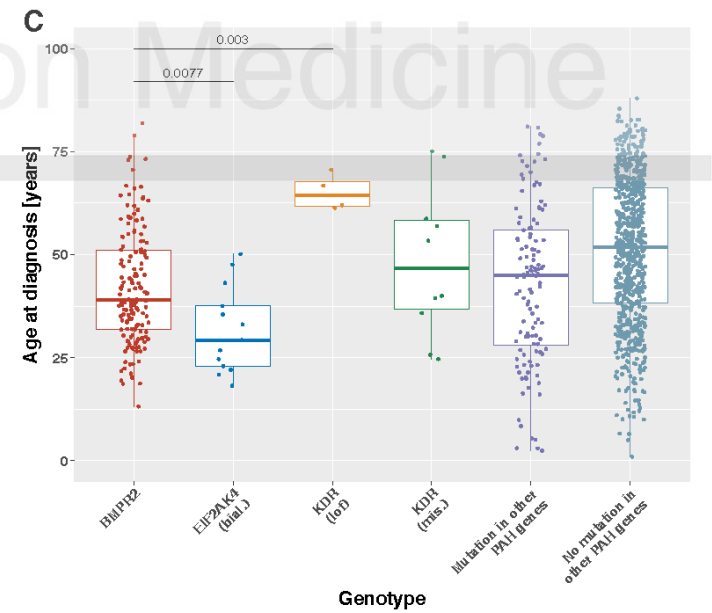
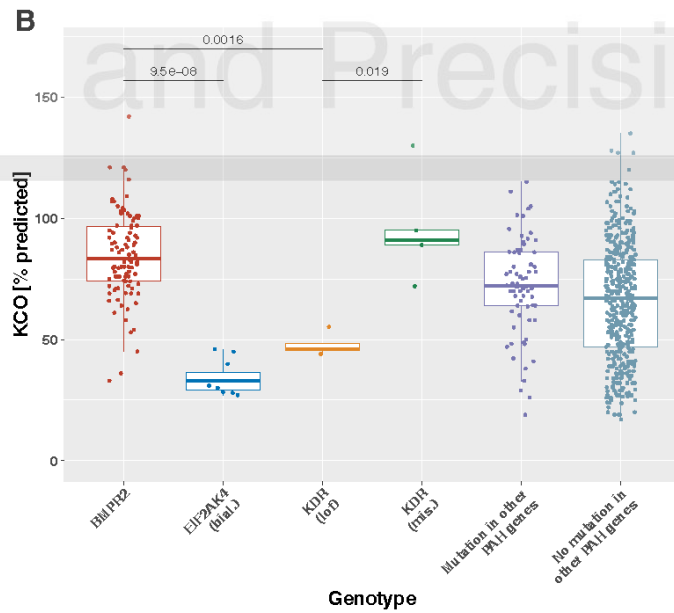
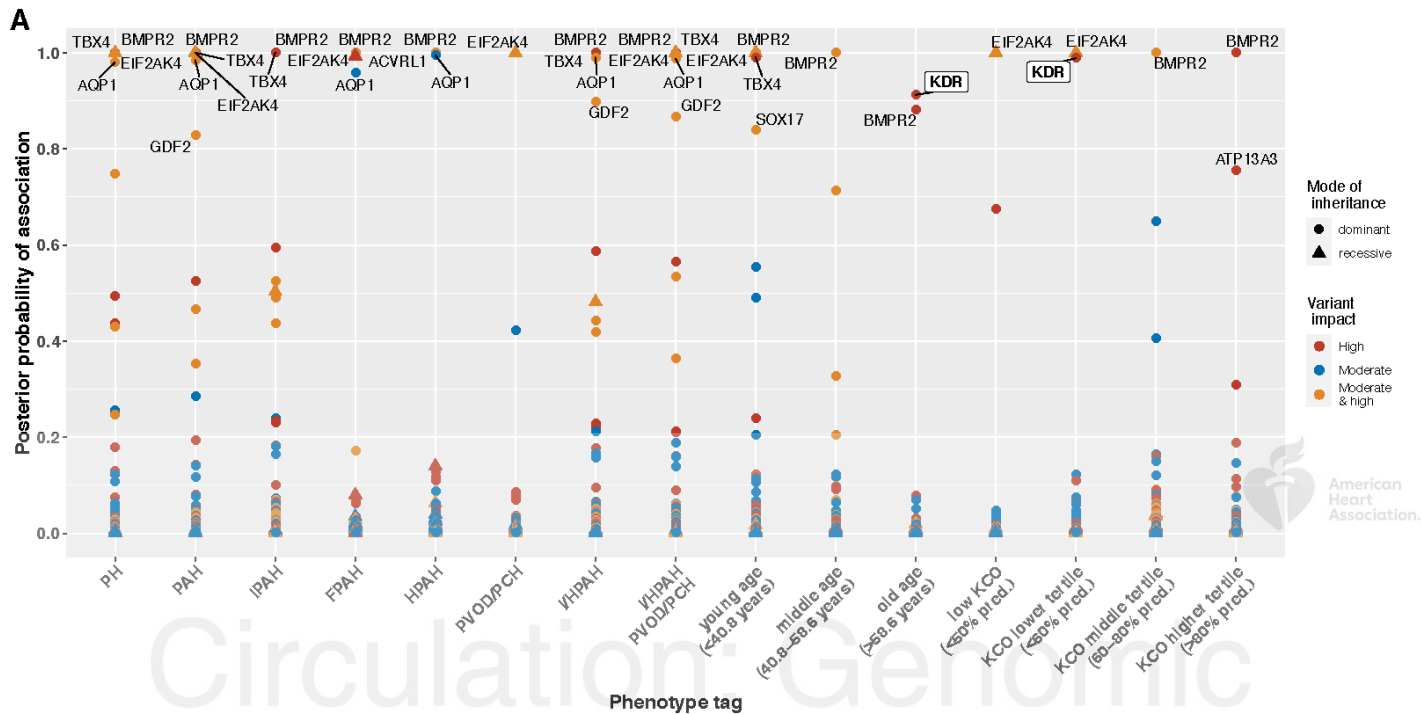
B**Cases**

KCO in the lower tertile
included

KCO in the middle or higher tertile
excluded



Missing KCO measurement
excluded

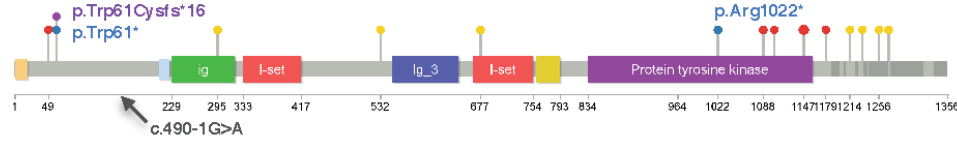


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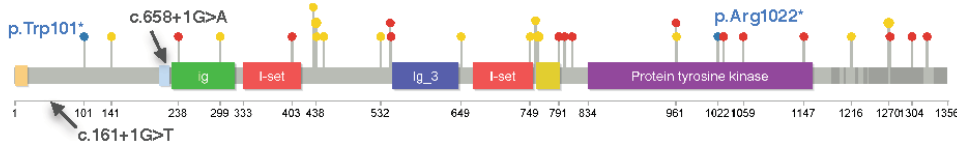
A

KDR SNVs and indels

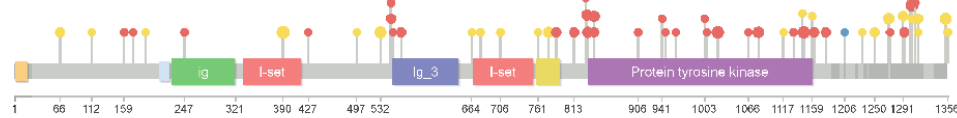
NBR PAH cases



US Biobank PAH cases



NBR non-PAH controls



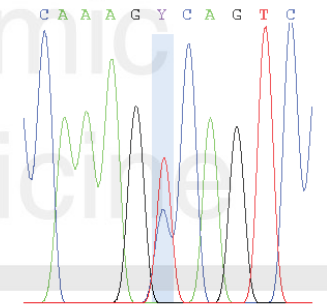
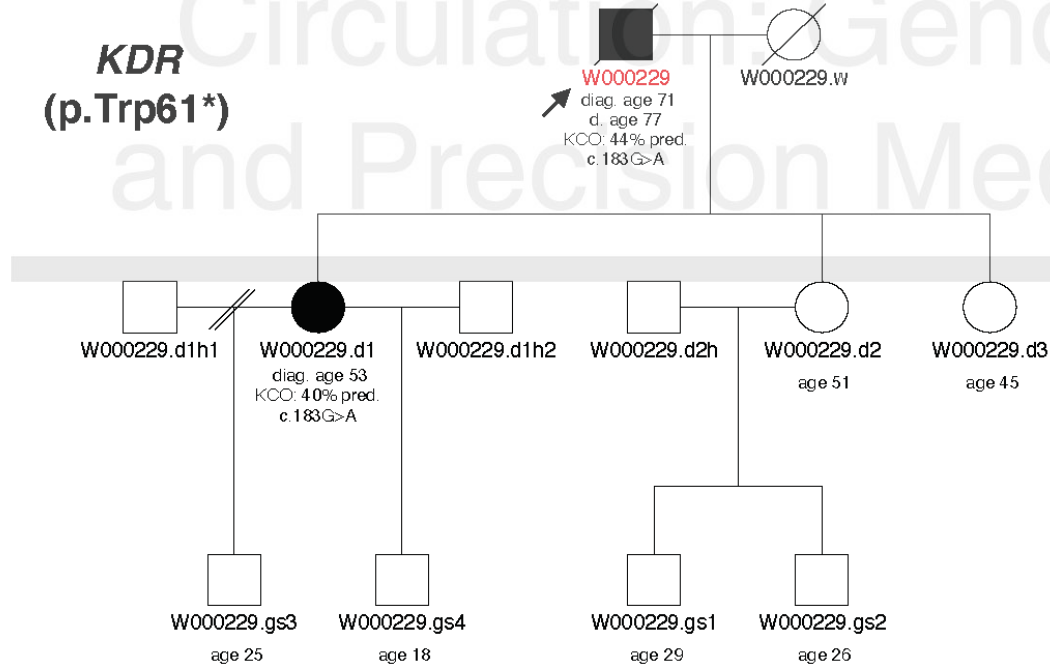
Consequence type

- Stop gained
- Frameshift
- Splice site
- Missense (SIFT / PolyPhen-2 agree)
- Missense (SIFT / PolyPhen-2 disagree)

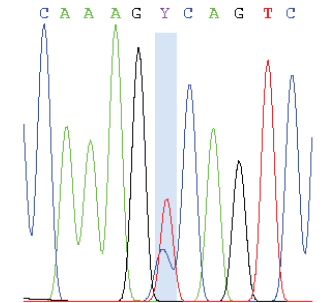
KDR variant consequence	NBR		USBB
	Cases	Controls	Cases
frameshift	1	0	0
nonsense	2	1	2
splice acceptor	1	0	0
splice donor	0	0	2
missense	13	107	31

B

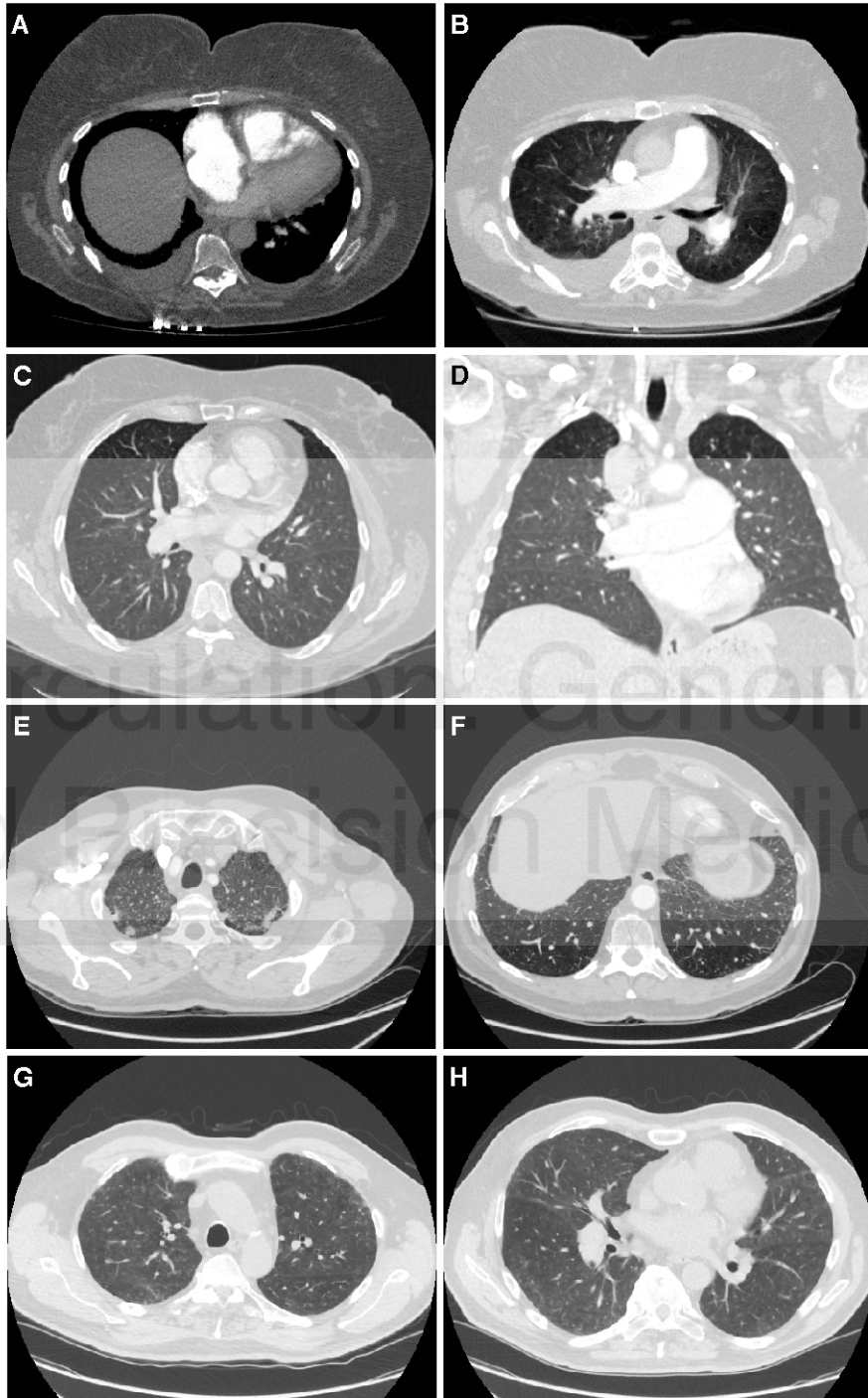
KDR (p.Trp61*)



W000229



W000229.d1



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