Characterisation of the Cullin-3 mutation that causes a severe form of familial hypertension and hyperkalaemia

Frances-Rose Schumacher, Keith Siew, Jinwei Zhang, Clare Johnson, Nicola Wood, Sarah E Cleary, Raya S Al Maskari, James T Ferryman, Iris Hardege, Yasmin, Nichola L Figg, Radoslav Enchev, Axel Knebel, Kevin M O'Shaughnessy & Thimo Kurz

Abstract

Deletion of exon 9 from Cullin-3 (CUL3, residues 403–459: CUL3<sup>403–459</sup>) causes pseudohypoaldosteronism type IIE (PHA2E), a severe form of familial hyperkalaemia and hypertension (FHHt). CUL3 binds the RING protein RBX1 and various substrate adaptors to form Cullin-RING-ubiquitin-ligase complexes. Bound to KLHL3, CUL3–RBX1 ubiquitylates WNK kinases, promoting their ubiquitin-mediated proteosomal degradation. Since WNK kinases activate Na/Cl co-transporters to promote salt retention, CUL3 regulates blood pressure. Mutations in both KLHL3 and WNK kinases cause PHA2 by disrupting Cullin-RING-ligase formation. We report here that the PHA2E mutant, CUL3<sup>403–459</sup>, is severely compromised in its ability to ubiquitylate WNKs, possibly due to altered structural flexibility. Instead, CUL3<sup>403–459</sup>-auto-ubiquitylates and loses interaction with two important Cullin regulators: the COP9-signalosome and CAND1. A novel knock-in mouse model of CUL3<sup>403–459</sup> closely recapitulates the human PHA2E phenotype. These mice also show changes in the arterial pulse waveform, suggesting a vascular contribution to their hypertension not reported in previous FHHt models. These findings may explain the severity of the FHHt phenotype caused by CUL3 mutations compared to those reported in KLHL3 or WNK kinases.

Keywords: cullin; CUL3; monogenic hypertension syndromes; proteasome; ubiquitin; WNK/SPAK/OSR pathway

Subject Categories: Genetics, Gene Therapy & Genetic Disease; Post-translational Modifications, Proteolysis & Proteomics; Urogenital System

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Introduction

Blood pressure is controlled in part by the WNK kinase pathway, which acts to modulate the activity of key cation-chloride co-transporters in the kidney (Kahle et al., 2008; Alessi et al., 2014). Considerable insight into this pathway has come from exploring the molecular basis for mutations described in patients with a rare monogenic form of hypertension called familial hypertension and hyperkalaemia [FHHt, also known as pseudohypoaldosteronism (PHA2); OMIM: 145260]. Mutations in the genes encoding WNK kinases have been shown to increase protein levels of WN1 and WNK4 in the kidney, leading to increased activation of this pathway (Wilson et al., 2001; Kahle et al., 2008). When activated by phosphorylation, the WNK kinases phosphorylate and activate the downstream kinases, SPAK (SPS1-related proline/alanine-rich kinase) and OSR1 (oxidative stress-responsive kinase 1), which in turn phosphorylate and activate cation-chloride co-transporters in the kidney, such as NCC (Na<sup>+</sup>/Cl<sup>−</sup> co-transporter), which leads to increased sodium ion reabsorption (Vitari et al., 2005; Delpire & Gagnon, 2008; Richardson et al., 2008, 2011). It is this up-regulation of the WNK signalling pathway that is responsible for the salt retention that drives the hypertension in FHHt. These patients also present with disturbed plasma electrolytes especially hyperkalaemia and a metabolic acidosis, partly driven by NCC activation (Boyden et al., 2012; Osawa et al., 2013; Tsuji et al., 2013; Glover et al., 2014; McCormick et al., 2014). As FHHt patients are salt-loaded through NCC activation, the hypertension and electrolyte disturbance respond to either salt restriction or low doses of thiazide diuretics (Gordon & Hodsman, 1986; Mayan et al., 2002).

More recently, the ubiquitin–proteasome system (UPS) was shown to have a critical upstream role in regulating blood pressure by modulating WNK kinase levels (Mori et al., 2013; Ohta et al.,...
2013; Wakabayashi et al., 2013; Wu & Peng, 2013). This was first suggested when patients with severe forms of FHHt were described with mutations in Cullin-3 (CUL3; PHA2E), and one of its substrate adaptors, KLHL3 (PHA2D) (Boyden et al., 2012; Louis-Dit-Picard et al., 2012; OMIM:145260). CUL3 belongs to the Cullin-RING ligase (CRL) family that form the largest class of ubiquitin E3 ligases in the cell. Cullins are elongated proteins, which confer an extended structure to active CRL complexes (Zheng et al., 2002; Duda et al., 2011). Through their C-terminus, Cullins constitutively bind to a small RING finger protein, either RBX1 or RBX2, which forms the catalytic core of the ligase complex. The binding to various substrate-adaptor proteins at the N-terminal domain enables CUL3-RBX1 to appropriately ubiquitylate a plethora of cellular substrates (Wee et al., 2005; Harper & Tan, 2012). In this way, CUL3 acts as a direct scaffold to direct ubiquitylation, as C-terminally bound RBX11 binds ubiquitin-conjugating E2 enzymes that are charged with ubiquitin, while simultaneously binding to the substrate adaptor, in this instance KLHL3. The overall structural rigidity of the Cullin, in particular between the elongated N-terminus and the globular C-terminus, is important for the ubiquitin E3 ligase function of the complex. Alterations in the connecting sequence between the Cullin N- and C-terminus are thought to change the relative positioning of the RING domain towards the bound substrate, which can prevent substrate modification (Zheng et al., 2002; Liu & Nussinov, 2011). Efficient CRL ubiquitylation also requires the modification of a conserved C-terminal lysine residue (Lys 721 in CUL3) by a single molecule of the ubiquitin-like protein Nedd8 (Pintard et al., 2003). This modification, auto-catalysed by CUL3-RBX1, is aided by Nedd8-E3 ligases (Kurz et al., 2005; Rabut et al., 2011) and alters the flexibility of the C-terminal Cullin-RBX1 module to activate E3 ligase activity (Duda et al., 2008; Scott et al., 2014). While neddylation is required for efficient ubiquitylation, studies show that the cycling between a neddylated and deneddylated state is crucial in a cellular context (Cope et al., 2002; Pintard et al., 2003). As such, Cullin interactions with the deneddylation complex COP9-signalsome (CSN), a metalloprotease, are also required for activity. Another regulator of CRLs, CAND1, interacts with the N- and C-terminus of Cullins only in their deneddylated state to promote the exchange of substrate adaptors (Liu et al., 2002; Goldenberg et al., 2004; Pierce et al., 2013; Wu et al., 2013; Zemla et al., 2013).

In the context of PHA2, KLHL3 is the critical adaptor that links the WNK kinases to CUL3-RBX1, enabling CUL3 to directly ubiquitylate the bound WNK kinase. In earlier work, we and others showed the KLHL3 mutations described in FHHt (PHA2D) patients disrupt the formation of the complex and prevent WNK ubiquitylation, leading to WNK stabilisation and increased flux through the WNK kinase pathway (Mori et al., 2013; Ohta et al., 2013; Shibata et al., 2013; Wakabayashi et al., 2013; Wu & Peng, 2013; Schumacher et al., 2014; Susa et al., 2014). Some patients with FHHt have mutations in WNK kinases, which promote WNK stabilisation by disrupting WNK-KLHL3 interactions (Wakabayashi et al., 2013; Schumacher et al., 2014) or, in some cases, by the increased expression of WNK mRNA (Wilson et al., 2001).

While FHHt patients with CUL3, KLHL3 or WNK mutations present with the same disease, those with mutations in CUL3 display a more severe phenotype, evident in terms of both an earlier age-of-onset and the degree of hypertension and electrolyte disturbance reported (Boyden et al., 2012; Osawa et al., 2013; Tsuji et al., 2013).

To date, all the reported CUL3 PHA2E mutations are heterozygous, transmit as autosomal dominant traits and result in the deletion of the amino acids encoded for by exon 9 of the CUL3 mRNA (residues 403–459) (Boyden et al., 2012; Grimm et al., 2012; Osawa et al., 2013; Tsuji et al., 2013; Glover et al., 2014). Nevertheless, precisely why and how this truncated form of CUL3 (hereafter referred to as CUL3403–459) causes a more severe phenotype is currently unclear. We have investigated this question using a biochemical in vitro approach to understand the molecular defects caused by the mutation and a mouse model of PHA2E to better understand the physiological basis of the effects of CUL3403–459. This has uncovered a number of critical molecular and physiological factors that enhance our understanding of Cullin-RING ligase mechanisms, the physiology of CUL3, and importantly its pathophysiology in FHHt patients.

**Results**

**CUL3403–459 forms an active Cullin-RING ligase complex**

To investigate the molecular defect of CUL3403–459, we determined whether this form of CUL3 was able to build a CRL complex. First, we established that CUL3403–459 binds RBX1 similar to CUL3WT, by showing that both versions of CUL3 form a stable complex with RBX1 when co-expressed in insect cells (Fig 1A). A second critical interaction for a functional CRL is the ability to bind substrate-adaptor proteins, and in the context of WNK kinase modification, CUL3 must interact with KLHL3 (Lamark & Johansen, 2012; Ji & Privé, 2013; Canning & Bullock, 2014). The 57-residue deletion in CUL3403–459 lies in a structurally distinct domain of CUL3 and would not be expected to perturb binding to substrate adaptors. We confirmed that CUL3403–459 indeed retains the ability to bind KLHL3 by in vitro pull-down assays with purified proteins (Fig 1B). We also showed this interaction is maintained in a cellular system, using ectopically expressed FLAG-tagged CUL3 to co-precipitate endogenous KLHL3 inFLAG immunoprecipitates (Fig 1H, lower panel). Thus, CUL3403–459 maintains crucial interactions important for ubiquitin-ligase function. Given this, we tested whether CUL3403–459-RBX1 also maintains the ability to hydrolyse the thioester bond between the catalytic cysteine residue of a recruited E2 enzyme and ubiquitin (E2–UB), a critical step in ubiquitylation and a measure for catalytic activity. Using ubiquitin-release assays from charged E2 enzymes, we show that CUL3403–459 hydrolyses E2–UB more efficiently than CUL3WT, demonstrating a functionally intact CUL3403–459-RBX1 catalytic core, while also suggesting potential hyper-activity (Fig 1C). Together, these results demonstrate that CUL3403–459 maintains interactions and basic functionality critical for CRL activity.

**Structural modelling suggests CUL3403–459 is more flexible than CUL3WT**

To better understand how the deletion in CUL3403–459 may affect the CRL, we predicted the structure of CUL3, using closely related Cullin-1 (CUL1; Fig 1D; Appendix Fig S1). The region deleted in CUL3403–459 likely forms a three-helical bundle juxtaposed to the globular C-terminal domain. Removal of these helices would fuse together two short unstructured regions to create a longer loop,
which, as well as shortening the elongated structure of the CRL complex, may increase the flexibility between the two domains. If correct, this model explains how RBX1 and KLHL3 binding is maintained (Munir et al., 2007; Calabrese et al., 2011), while also suggesting the function of the ligase may be compromised due to the altered relative orientation of the CRL components towards each other. Studies investigating the structure of CUL1 demonstrated the importance of a rigid Cullin conformation for substrate ubiquitylation (Zheng et al., 2002; Appendix Fig S1). It is therefore possible that due to the presence of a predicted unstructured flexible loop, CUL3\(^{403-459}\) may similarly sample non-productive conformations.

CUL3\(^{403-459}\) has altered Nedd8-ligase activity and is unable to interact with the CRL regulators COP9-signalsome and CAND1

Available structural data of CRL proteins bound to Cullin regulators suggest the deleted region in CUL3\(^{403-459}\) may also influence neddylation and binding of the CSN (Lingaraju et al., 2014) and of CAND1 (Goldenberg et al., 2004; Fig EV1). This prompted us to examine whether these interactions are affected by the PHA2E mutation.

To investigate the ability of CUL3\(^{403-459}\) to be auto-modified with Nedd8, we used in vitro neddylation assays with purified CUL3, Nedd8 E1 (APPBP1-UBA3), Nedd8 E2 (UBE2M) and Nedd8. As Nedd8 forms a covalent isopeptide bond with CUL3, neddylation can be visualised as a slower-migrating band on SDS-PAGE. We found CUL3\(^{403-459}\) retains the ability to auto-neddylate (Fig 1E, Appendix Fig S2A). However, CUL3\(^{403-459}\) appears less efficient at transferring Nedd8, as unlike for CUL3\(^{WT}\), some unmodified CUL3\(^{403-459}\) remained even 45 min after neddylation was initiated. Moreover, while CUL3\(^{WT}\) was only mono-neddylated, multiple Nedd8 molecules were attached to CUL3\(^{403-459}\) (Fig 1E). Both of these differences in neddylation are consistent with increased structural flexibility of CUL3\(^{403-459}\).

As reported (McCormick et al., 2014), we also noticed that CUL3\(^{403-459}\) neddylation is elevated in cells (Fig 1F). CRLs cycle between neddylated and de-neddylated states to maintain activity (Pintard et al., 2003). Therefore, the increased cellular neddylation of CUL3\(^{403-459}\) could reflect a decrease in de-neddylation by the isopeptidase CSN. The structure of the CSN bound to
CUL1 supports this idea, as the Cullin C-terminal domain, including the region equivalent to residue 403–459 in CUL3, contacts the CSN subunits directly (Lingaraju et al., 2014). To investigate this possibility, we blocked neddylation using an inhibitor of the Nedd8-El enzyme (MLN4924; Millennium Pharmaceuticals) and monitored CUL3 de-neddylation over time. Consistent with decreased rates of deneddylation, CUL3
\(^{403–459}\) remained neddylated for longer than CUL3 WT (Fig 1G) and also showed markedly reduced binding to CSN subunits (Fig 1H, Appendix Fig S2B). As such, the observed elevated Neddd8 modification of CUL3
\(^{403–459}\) in cells is likely due to a combination of Nedd8 ligation to non-physiological lysine residues and of reduced CUL3 de-neddylation.

CRLs must be de-neddylated for a productive interaction with the substrate-adaptor exchange factor CAND1. Furthermore, the structure of CUL1 bound to CAND1 suggests that, regardless of neddylation state, CUL3
\(^{403–459}\) may be unable to form the required critical interactions (Goldenberg et al., 2004; Pierce et al., 2013; Wu et al., 2013; Zemla et al., 2013). We therefore investigated whether CUL3
\(^{403–459}\) was indeed defective in binding to CAND1. We treated cells expressing either FLAG-CUL3 WT or FLAG- CUL3
\(^{403–459}\) with the neddylation inhibitor MLN4924 to force all of CUL3 into the non-neddylated state and then determined interaction with CAND1 by immunoprecipitation. While CUL3 WT strongly bound to CAND1 in these assays, CUL3
\(^{403–459}\) was unable to interact with CAND1 (Fig 2H, Appendix Fig S2B). We confirmed these findings using in vitro assays with purified proteins (Fig 1I). These data advocate that structural perturbations prevent CUL3
\(^{403–459}\) from binding to CAND1. In combination with the impairment of CSN binding, such defects suggest that the regulation of CUL3
\(^{403–459}\) in cells is likely severely disrupted.

**CUL3
\(^{403–459}\) displays enhanced auto-ubiquitylation**

Our data indicate that the regulation of CUL3
\(^{403–459}\) is defective in cells, and we wanted to test whether its ubiquitylation activity is similarly affected. McCormick et al suggested that CUL3
\(^{403–459}\) may stabilise WNK kinases by ectopically degrading the substrate adaptor KLHL3, preventing the formation of functional CUL3-KLHL3 complexes (McCormick et al., 2014). Some CRLs are known to ubiquitylate bound substrate adaptors (Wee et al., 2005), and it is feasible that altering the CUL3 backbone structure could indeed lead to erroneous ubiquitylation of KLHL3. In our in vitro system, we confirmed that CUL3
\(^{403–459}\) directly modifies KLHL3 with a greater efficiency than CUL3 WT (Fig 2A, Appendix Fig S2C). Interestingly, we observed that CUL3
\(^{403–459}\) also exhibited markedly increased auto-ubiquitylation (Fig 2B, Appendix Fig S2D).

Auto-ubiquitylation is often utilised as a read-out for ubiquitin-ligase activity. In a cellular context, self-modification can lead to auto-degradation of the ligase (Silke et al., 2005). If CUL3
\(^{403–459}\)
Figure 2.
Cullin-3 and defective WNK ubiquitylation in PHA2E
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D 2006). In these assays, CUL3

et al had increased to five, all of which were C-terminal to the original

ubiquitylation towards a bound WNK substrate in in vitro ubiquitylation

assays using purified components. As we previously showed,

lysine residues by analysing which lysine residue(s) carry di-glycine

lysine residues. (C) Activity assays contain methylated ubiquitin, a form of ubiquitin incapable of

forming ubiquitin chains; as such, higher band shifts reflect the attachment of mono-ubiquitin to one more lysine residue on CUL3 WT or CUL3 403–459, respectively. (D) Activity assay performed as in (C) with methyl-ubiquitin, the boxes shown on the gel are indicative of the gel pieces excised for mass spectrometry analysis in (F).

E Schematic representation (to linear residue scale) highlighting the domains of CUL3 with schematic representation of the lysine residues of CUL3 WT or CUL3 403–459, modified at 5 and 45 min, respectively, in the in vitro ubiquitylation assay shown in (D) and as determined by mass spectrometry.

F Structural docking model of CUL3 403–459 based on the structure of full-length CUL1 (1LDK) using Chimera (see Materials and Methods). The NTD is coloured green, while the CTD is red, and the four panels sample possible positioning of the CTD relative to the NTD in CUL3 403–459, assuming full flexibility of the linker after deletion of three helices encoded by exon 9.

Source data are available online for this figure.

CUL3 403–459 displays increased auto-ubiquitylation and ubiquitylation of KLHL3.

A–D In vitro ubiquitylation assays with purified proteins. As ubiquitin is covalently attached to the substrate lysine residue, the appearance of higher molecular weight protein bands reflects the modification of the protein with ubiquitin. All assays contain purified UB1, UB2D3, ubiquitin, 0.1 mM ATP, 1 mM MgCl2, and were buffered in 50 mM HEPES, 150 mM NaCl and incubated at 30°C for the time indicated. Reactions were stopped by the addition of SDS–PAGE, staining with Coomassie blue, or detection with the indicated antibody following immunoblotting enabled the visualisation of ubiquitylation. (A) to determine the relative modification of KLHL3 by CUL3 WT and CUL3 403–459, KLHL3 was included into the reaction at 2 μmol concentration over CUL3 WT and CUL3 403–459. (B, C) Reactions serve to determine basal auto-ubiquitylation of CUL3 WT or CUL3 403–459 and do not contain KLHL3 or other potential substrate proteins. Lysine residues on CUL3 WT or CUL3 403–459 act as the substrate. (B) High molecular weight bands reflect ubiquitin chain linkages on CUL3 or the multiple mono-ubiquitylation of a number of CUL3 lysine residues. (C) Activity assays contain methylated ubiquitin, a form of ubiquitin incapable of forming ubiquitin chains; as such, higher band shifts reflect the attachment of mono-ubiquitin to one more lysine residue on CUL3 WT or CUL3 403–459.

E Structural docking model of CUL3 403–459 using Chimera (see Materials and Methods). The NTD is coloured green, while the CTD is red, and the four panels sample possible positioning of the CTD relative to the NTD in CUL3 403–459, assuming full flexibility of the linker after deletion of three helices encoded by exon 9.

Source data are available online for this figure.

CUL3 403–459 modifies lysine residues inaccessible to CUL3 WT, this could explain the increased auto-ubiquitylation. Alternatively, CUL3 403–459 could build longer ubiquitin chains on the same lysine residues that are also targeted by CUL3 WT. To explore these possibilities, we performed in vitro ubiquitylation assays with methylated ubiquitin, a form of ubiquitin unable to build ubiquitin chains (Kirisako et al, 2006). In these assays, CUL3 403–459 was auto-modified on multiple lysine residues, while CUL3 WT only mono-ubiquitylated one residue (Fig 2C). Therefore, the observed increase in auto-ubiquitylation is most likely due to erroneous modification of multiple lysine residues on CUL3 403–459, which is consistent with an increase in structural flexibility as suggested by our model. Alternatively, the mutation may change the active site environment of CUL3, exposing more lysine residues close to the E2–UB binding site. To distinguish between these possibilities, we mapped the lysine residues modified in our in vitro assay system using mass spectrometry. We performed in vitro auto-ubiquitylation reactions of CUL3 with methylated ubiquitin for either 5 or 45 min and determined the ubiquitylated lysine residues by analysing which lysine residue(s) carry di-glycine remnants after trypsin digest (Fig 2D and E; Kim et al, 2011). Using this approach, we found that after 5 min, CUL3 WT was only auto-ubiquitylated on a single lysine residue (K336), while ten lysine residues on CUL3 403–459 were modified (Fig 2D and E). After running the reaction for 45 min, the modification sites on CUL3 WT increased to five, all of which were C-terminal to the original site and contained within the globular C-terminal Cullin domain. The ubiquitylation sites on CUL3 403–459, however, had increased to 25 sites distributed along the entire length of the protein (most N-terminal: K47; most C-terminal: K743). These results support our model that the CUL3 deletion induces increased structural flexibility, leading to the erroneous auto-ubiquitylation of sites normally not accessible to CUL3 WT (Figs 2E and F and EV2).

CUL3 403–459 cannot ubiquitylate WNK kinases

As CUL3 403–459 is able to more efficiently auto-ubiquitylate itself and KLHL3, we next asked whether it similarly shows increased ubiquitylation towards a bound WNK substrate in in vitro ubiquitylation assays using purified components. As we previously showed, CUL3 WT efficiently ubiquitylates WNK kinases in vitro (Fig 3; Ohta et al, 2013). CUL3 403–459, however, while maintaining the ability to efficiently modify itself and KLHL3, was not able to modify either WNK1 or WNK4 (Fig 3A and B). As described for CUL1 (Zheng et al, 2002), increasing the flexibility of the Cullin backbone can lead to a decrease in substrate modification, while promoting excessive modification of other CRL components (Zheng et al, 2002). Thus, increased structural flexibility of CUL3 403–459 may lead to similar defects. If so, then it becomes important to understand how this in vitro finding can explain the increased WNK stabilisation found in vivo. It is possible that only one of the three defects, the lack of substrate ubiquitylation, the increased Cullin auto-ubiquitylation or the increased KLHL3 ubiquitylation, is the major driver for the observed phenotypes. Decisively, KLHL3 levels are not decreased in the kidney of CUL3 WT/A403–459 mice, while the overall CUL3 levels are lower. This suggests that KLHL3 degradation does not contribute to the observed PHA2E phenotype, and instead, we propose that CUL3 403–459 stabilises WNK levels in two ways: firstly, CUL3 403–459 exquisitely auto-ubiquitylates, leading to its own degradation; and secondly, any CUL3 403–459 protein that escapes degradation cannot promote the ubiquitylation of the WNK kinases. Importantly, all PHA2E patients are heterozygous for the mutation, suggesting that with respect to blood pressure regulation either CUL3 403–459 behaves as a dominant-negative or CUL3 WT is haploinsufficient. Our data demonstrate that the mutation causes a loss-of-function, so a dominant-negative effect seems less likely. We also find that in vitro, equimolar amounts of CUL3 403–459 do not inhibit CUL3 WT (Fig 3C), further supporting the concept of haploinsufficiency.

CUL3 WT/A403–459 mice have increased flux through the WNK kinase pathway and typical FHHT phenotypic features of electrolyte imbalance

To complement our in vitro findings and to determine more accurately what occurs in PHA2E patients, we engineered a knock-in (KI) mouse carrying the same exon 9 deletion of CUL3 as reported in PHA2E pedigrees (CUL3 403–459; Appendix Fig S3). Despite setting up heterozygous mating pairs, no homozygous KI mice were born suggesting that, like deletion of the entire coding region of CUL3, the homozygous deletion of residues 403–459 is lethal (Singer et al, 1999). Nevertheless, all the reported PHA2E pedigrees are heterozygous, so we phenotyped the heterozygous KI mice. Utilising both immunoblotting of whole kidney lysates and immunofluorescence

Figure 2. CUL3 403–459 displays increased auto-ubiquitylation and ubiquitylation of KLHL3.
confocal microscopy of kidney sections, we determined the relative abundance and distribution of proteins involved in the WNK signalling pathway. Whole kidney lysate was analysed by immunoblotting confirming the expression of CUL3 and the presence of an additional band corresponding to the expected PHA2E form, CUL3\(^{3403-459}\), in the heterozygous mice (Fig 4A). Increased levels of WNK4 and the downstream components of the WNK kinase pathway, reported in other mouse models of FHHt, were also observed (Fig 4A). Specifically, there were striking increases in both total and phosphorylated forms of SPAK and NCC (Fig 4A). We also confirmed the expression of KLHL3 and CUL3 in normal human kidney lysates (Fig 4B). In other mouse models of FHHt, were also observed (Fig 4A). Specifically, there were striking increases in both total and phosphorylated forms of SPAK and NCC (Fig 4A). We also confirmed the expression of KLHL3 and CUL3 in normal human kidney lysates (Fig 4B). In contrast, CUL3\(^{3403-459}\) appears to be less abundant than CUL3\(^{WT}\) in the heterozygous mice. These observations suggest that CUL3\(^{3403-459}\) does not promote KLHL3 degradation in vitro and, moreover, that the lower levels of CUL3\(^{3403-459}\) may be due to the propensity of CUL3\(^{3403-459}\) to self-modify presumably causing degradation through the UPS. The lower level of observable CUL3\(^{3403-459}\) may also be due to non-proteolytic hyper-modification of CUL3\(^{3403-459}\) with either Nedd8 or ubiquitin. Indeed, this may in part be the case, as incubation of kidney extract at room temperature to promote de-conjugation of Nedd8 or ubiquitin increased the abundance of the faster migrating CUL3\(^{3403-459}\) band, but did not restore it to CUL3\(^{WT}\) levels (Fig 4A, lower panel). Regardless of precisely why, the stabilisation of WNK4 and increased signalling through the WNK cascade is consistent with a PHA2E model.

Over-activation of the WNK pathway would be expected to perturb electrolyte homeostasis. We confirmed this by measuring urine electrolytes, blood biochemistries and aldosterone levels in the mice (Figs 4C and D and EV3). The CUL3\(^{WT/Δ403-459}\) mice showed the typical electrolyte disturbance of FHHt with hyperkalaemia, hyperchloraemia and a compensated metabolic acidosis, with the pH maintained by an increased respiratory drive to increase
Figure 4.

A

Mouse kidney extract

WT/WT  WT/Δ403-459

Shorter exposure

WT/WT  WT/Δ403-459

CUL3  CUL3Δ

KLHL3

WNK4

-SPAK Thr373-P

-SPAK total

-NCC Thr60-P

-NCC total

-Actin

IP:CUL3

WT/WT  WT/Δ403-459

IB: CUL3

B

Human kidney extract

65

KLHL3

42

β-actin

42

β-actin

C

Plasma aldosterone (pg/ml)

n = 22

n = 18

D

Plasma level (mmol/L)

K+  CI-

mMol/L

pH

BE

KPa

pH  HCO3-

BE  pCO2

CUL3^WT

(n = 13)

CUL3^WT/Δ403-459

(n = 13)
CO₂ removal and hence reduced total CO₂ and pCO₂ levels. The mice also had elevated aldosterone levels (Fig 4C) driven by the hyperkalaemia without pronounced hypercalcaemia as evidenced by similar haematocrict levels (Fig EV3). Hypermagnesaemia and hyperphosphataemia were noted on detailed electrolyte analysis of plasma and urine, which are the converse of the abnormalities reported in Gitelman syndrome (Fig EV3; Akhtar & Hafeez, 2009; Rafiqi et al, 2010; Pathare et al, 2012; Ali et al, 2013; Zhang et al, 2015). Additionally, we also found that the CUL3WT/403–459 mice have significantly lower body weight and body length relative to the WT mice (Appendix Fig S4). This is consistent with Gordon’s original observation that children with PHA2 may present with a low percentile body weight and height for their age (Gordon, 1986) and more recent reports that CUL3WT/403–459 patients are growth retarded (Osawa et al, 2013; Tsuji et al, 2013).

**WNK4 and SPAK accumulate and form puncta in the distal convoluted tubule of CUL3WT/403–459 mice**

Using immunofluorescence confocal microscopy of kidney sections, we studied the distribution of proteins involved in the WNK kinase cascade in both CUL3WT/WT and CUL3WT/403–459 mice (Fig 5A). CUL3WT/403–459 mice showed a similar CUL3 distribution compared to both the WT mice, and to the WT human kidney (Fig 5A and B). This was in keeping with the abundance observed in the kidney immunoblots (Fig 4A). In contrast, staining for WNK4 and SPAK in the CUL3WT/403–459 mice revealed these proteins to have a striking punctate appearance in the DCT, but not in adjacent segments of the nephron including the thick ascending limb (TAL; Figs 5A and EV4). While WNK1 and OSR1 puncta have been reported in the DCT of SPAK knockout mice, these were smaller in size and far less numerous (Grimm et al, 2012). As these puncta do not co-localise with the lysosomal marker LAMP1 or form ubiquitin-containing aggregrates (Figs 5A and EV4), they may represent an accumulation of WNK and SPAK proteins within a protein-processing compartment, such as the trans-Golgi network or activation of secondary protein degradation pathways or storage compartments, due to failed proteosmal clearance (D’Urso et al, 1998; Lamark & Johansen, 2012; Wolff et al, 2014).

**CUL3WT/403–459 mice have hypertension and a novel vascular phenotype**

Over-activation of the WNK pathway as observed in our CUL3WT/403–459 mice (Figs 4 and 5) should lead to salt retention and elevated blood pressure. We therefore measured the blood pressure (BP) of the CUL3WT/403–459 mice using a carotid pressure transducer catheter under general anaesthesia. Both male and female CUL3WT/403–459 mice exhibited significantly higher BP relative to CUL3WT/WT littermates (Fig 6A).

Strikingly, we also observed a previously unreported haemodynamic change in the blood pressure trace of the CUL3WT/403–459 mice. The systolic aortic pressure wave of the heterozygous mice is augmented, while the diastolic relaxation (τ) is significantly slowed (Munir et al, 2007; Fig 6B). These changes suggest early wave reflection of the pressure wave and compliance changes consistent with a stiffened vascular tree. This stiffening could reflect a primary change in the contractile state of vascular smooth muscle, so to explore this we studied the in vivo vasoconstrictor responses to intravenous administration of phenylephrine and angiotensin II (Fig 6C; Bergaya et al, 2011). The resulting dose–response curves showed that the maximal constrictor response to both vasopressor agents was substantially higher in the CUL3WT/403–459 mice.

**CUL3 and KLHL3 are expressed in the mouse and human aorta, and CUL3WT/403–459 mice have aortic wall thickening**

We also looked for evidence of biochemical changes in the vasculature and confirmed the expression of CUL3 and KLHL3 in mouse and human aorta by immunoblot analysis (Fig 7A and B, Appendix Fig S5). Similar to the kidneys (Fig 4A), no differences were seen in KLHL3 expression between CUL3WT/403–459 and CUL3WT mice, while the levels of CUL3 WT protein were lower in CUL3WT/403–459 aorta. To address the altered contractility in the vessel wall in vivo, we also measured the level of pMYPT1 in aortic vessel lysates and showed a significant 1.7-fold increase (Feng et al, 1999; Somlyo & Somlyo, 2003; Appendix Fig S6). Image analysis of mouse and human aortae confirmed that both have strong CUL3 staining of the vascular endothelium and smooth muscle cells of the intimal and medial layers of the vessel wall (Fig 7C and D). Finally, to explore...
Figure 5.

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Cullin-3 and defective WNK ubiquitylation in PHA2E

Frances-Rose Schumacher et al

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further the vascular phenotype, we performed a morphometric analysis of the aortae of CUL3 mice that confirmed significant thickening of the vessel wall in the heterozygote mice (Fig 7E). This probably reflects hyperplasia/hypertrophy of the vascular smooth muscle cells in the medial layer, as the numbers of elastic laminae were not significantly different (Fig 7E).

Discussion

This study describes the first successful PHA2E mouse model involving CUL3 and highlights the molecular differences and defects of the mutant CUL3 protein. The deletion of 57 amino acids in CUL3\(^{403-459}\) does not affect the ability of the bound RING domain to hydrolyse E2-UB, and as such CUL3\(^{403-459}\) is still an active E3 ligase. Instead, our structural predictions and experimental findings suggest that the CUL3\(^{403-459}\) CRL scaffold lacks a level of rigidity required for substrate ubiquitylation. Our results are consistent with a previous report that demonstrated ubiquitylation of the Cullin-1 substrate (p27) was abrogated when a linker was inserted between the NTD and CTD of the closely related CUL1 protein (Zheng et al., 2002). Thus, the CUL3\(^{403-459}\) mutation is a novel physiological example of the importance of this CRL rigidity and to our knowledge the first example of a human mutation that impedes the scaffolding function of a Cullin. Our data are consistent with the working hypothesis that the CUL3\(^{403-459}\) mutation has increased structural flexibility. The deletion of three helices fuses together two unrestructured regions, which likely allows greater movement between the N-terminal and C-terminal domains of the Cullin. This prevents the complex from successfully directing ubiquitin towards a bound substrate, and instead, leads to increased auto-ubiquitylation of CUL3 and KLHL3. Importantly, it appears as if in vivo this auto-ubiquitylation triggers degradation of the mutant form of the Cullin, as only relatively low amounts of CUL3 ubiquitylation triggers degradation of the mutant form of the Cullin.

inhibit CUL3\(^{WT}\) or is haploinsufficient in the context of blood pressure regulation with the single functional copy of CUL3 unable to sustain the physiological need for WNK ubiquitylation. Our data support a haploinsufficiency model, as a large proportion of the mutant protein is removed from the cell by auto-degradation and in vitro CUL3\(^{403-459}\) does not inhibit CUL3\(^{WT}\). However, it is possible that in vivo the remaining CUL3\(^{403-459}\) may still sequester KLHL3 from CUL3\(^{WT}\) in a dominant-negative fashion, as in cells we observed increased binding between CUL3\(^{403-459}\) and KLHL3 (Fig 1H) and the reduced affinity for CAND1 may prevent the release of KLHL3 from non-functional CUL3\(^{403-459}\) complexes. These two possibilities could be addressed in a heterozygous mouse model carrying a full-length CUL3 deletion; under the haploinsufficiency model, it should show WNK stabilisation.

The CUL3\(^{WT}\)/CUL3\(^{403-459}\) mice have high blood pressure and up-regulated signalling in the context of the WNK kinase pathway, which parallels the WNK4 (DS68E; Yang et al., 2007; Chowdhury et al., 2013) and KLHL3 (RS28H; Mori et al., 2013) PHA2E mouse models previously described. Mechanistically, our data suggest CUL3\(^{403-459}\) promotes its own degradation, as it has a heightened propensity to self-ubiquitylate and is less abundant in mouse kidney extract. Hence, we would predict that PHA2E CUL3\(^{WT}\)/CUL3\(^{403-459}\) patients would similarly have lower overall CUL3 levels. As CUL3 ubiquitylates a number of other proteins, it seems remarkable then that PHA2E patients do not have other phenotypic manifestations. However, this may not be surprising if considered in the context of a haploinsufficiency model. BP homoestasis requires reactive and precise control of ion fluxes in the distal nephron, especially the DCT, as changes in intake and output of salt are constantly balanced. The phosphorylation of the NCC/NKCC ion transporters by WNK kinases appears to be regulated by two complementary systems: the regulation of total WNK protein levels by CUL3-KLHL3, and the level of WNK activation by phosphorylation. Currently, the sensing mechanisms that lead to WNK phosphorylation or the degradation of the WNK kinases by CUL3-KLHL3 are largely unknown. However, sufficient amounts of CUL3 must be available for signalling-dependent ubiquitylation of the WNK kinases. It is possible that having only half the amount of functional CUL3 within kidney cells is not enough for the system to respond rapidly. The presence in our knock-in mice of WNK4 puncta in vivo also suggests that DCT cells may deploy secondary protein degradation pathways to manage excess WNK proteins, such as the autophagy–lysosomal system. However, the nuclear proximity of the large puncta and the absence of lysosomal staining or formation of ubiquitylated aggregates suggest that these excess WNK proteins are shunted to protein quality control compartments such as the aggresome or
Figure 6.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>CUL3WT</th>
<th>CUL3WT/Δ403-459</th>
<th>Parameter</th>
<th>CUL3WT</th>
<th>CUL3WT/Δ403-459</th>
</tr>
</thead>
<tbody>
<tr>
<td>MAP (mmHg)</td>
<td>81.0 ± 0.7</td>
<td>93.9 ± 1.2 ***</td>
<td>PP (mmHg)</td>
<td>30.4 ± 0.6</td>
<td>38.3 ± 0.7 ***</td>
</tr>
<tr>
<td></td>
<td>[n = 18]</td>
<td>[n = 18]</td>
<td></td>
<td>[n = 18]</td>
<td>[n = 18]</td>
</tr>
<tr>
<td>DNP (mmHg)</td>
<td>92.0 ± 0.8</td>
<td>109.2 ± 1.3 ***</td>
<td>AP (mmHg)</td>
<td>13.0 ± 0.7</td>
<td>19.4 ± 0.8 ***</td>
</tr>
<tr>
<td></td>
<td>[n = 18]</td>
<td>[n = 18]</td>
<td></td>
<td>[n = 18]</td>
<td>[n = 18]</td>
</tr>
<tr>
<td>( \tau )</td>
<td>0.59 ± 0.01</td>
<td>0.65 ± 0.01 **</td>
<td>A1x (%)</td>
<td>42.6 ± 1.8</td>
<td>50.2 ± 1.6 **</td>
</tr>
<tr>
<td></td>
<td>[n = 14]</td>
<td>[n = 14]</td>
<td></td>
<td>[n = 18]</td>
<td>[n = 18]</td>
</tr>
</tbody>
</table>
Figure 6. Increased arterial stiffness contributes towards the hypertension of CUL3WT/A403–459 mice.

A Continuous blood pressure measurements were taken in mice under general anaesthesia with thermostatically controlled body temperature, right carotid artery pressure transducer catheterisation and ECG lead II heart rate monitoring. CUL3WT/A403–459 mice have elevated systolic (SBP) (**P = 1 × 10−12) and diastolic (DBP) blood pressure (**P = 8.5 × 10−5), although they also present with 7% lower heart rates (**P = 0.0485) when compared to CUL3WT mice. Two-tailed unpaired Student’s t-test; data are mean ± SEM.

B Pulse waveform analysis of blood pressure traces obtained in (A) reveals that CUL3WT/A403–459 have an increased pulse pressure (PP) [SBP – DBP] (**P = 2.1 × 10−12), augmentation pressure [AP] [SBP – anacrotic notch (AN) pressure] (**P = 35 × 10−7), dicrotic notch (DN) pressure (**P = 4.7 × 10−12) and mean arterial pressure (MAP) [1/3 × SBP + 2/3 × DBP] (**P = 1.3 × 10−6). This hypertensive phenotype is in part due to changes in vascular contractility in CUL3WT/A403–459 mice as evidenced by their higher augmentation index (AIX) [AP/PP] (**P = 0.0096), a marker of arterial stiffness, and is further supported by an increase in their diastolic pressure decay time constant (TE[0–97]) of 13.1 slope of diastolic pressure decay, measured 30 ms after DN and 20 ms before end DBP (**P = 0.0083), a surrogate marker of increased vascular resistance. Two-tailed unpaired Student’s t-test; data are mean ± SEM.

C In vivo dose–responses to phenylephrine and angiotensin II. After baseline measurements were obtained from mice in (A), the right external jugular vein was cannulated for administration of bolus doses of vasopressors in increasing half-log steps. Dose–response curves were generated and data analysed using a logistical function to determine the estimated dose producing half-maximal response (ED50) and maximum response (Emax). The fitted Emax for phenylephrine was increased in CUL3WT/A403–459 versus CUL3WT (183.9 ± 25 versus 164.9 ± 1.4 mmHg) (**P = 1 × 10−5), indicating an increased vasoconstrictor response to adrenergic stimulation elevating systolic blood pressure substantially above the CUL3WT maximum. Similarly, the fitted Emax for angiotensin II stimulation was higher in CUL3WT/A403–459 versus CUL3WT (155 ± 1.8 versus 138 ± 2.2 mmHg) (**P = 1.7 × 10−3). However, there was no change between CUL3WT/A403–459 versus CUL3WT sensitivity (as measured by ED50) to phenylephrine (36.6 ± 5.1 versus 31.4 ± 2.8 μg/kg bw) (P = 0.3778) or angiotensin II (0.77 ± 0.16 versus 0.93 ± 0.13 μg/kg bw) (P = 0.4403). Two-tailed unpaired Student’s t-test; data are mean ± SEM.

Source data are available online for this figure.

juxta-nucleolar quality control compartment (JUNQ) in an attempt to compensate for loss of proteasomal degradation (Lamark & Johansen, 2012; Wolf et al., 2014). Hence, while other CUL3 substrates may be unaffected in the CUL3WT/A403–459 mice, the rapid and signal-dependent switching of WNK ubiquitylation may be executed abnormally in these mice and PHA2E patients.

Mutations in CUL3, KLHL3, WNK1 and WNK4 are known to cause FHH with constitutive activation of the NCC co-transporter (Boyden et al., 2012; Osawa et al., 2013; Tsuji et al., 2013; Alessi et al., 2014; Glover et al., 2014). What has been less clear is why the mutation of CUL3 has resulted in a more severe form of FHH (PHA2E). Our unexpected discovery of a vascular phenotype in the CUL3WT/A403–459 mice leads us to speculate that their hypertension may not be driven by salt retention in the DCT alone. It is also worth noting that humans with hypertension (Kaess et al., 2012), but not previously in a hypertensive mouse model. It is also worth noting that augmentation of the aortic pressure wave is often not detected by brachial cuff BP measurements, so it could have gone unnoticed in a hypertensive mouse model. It is also worth noting that humans with hypertension (Kaess et al., 2012), but not previously in a hypertensive mouse model. It is also worth noting that augmentation of the aortic pressure wave is often not detected by brachial cuff BP measurements, so it could have gone unnoticed in a hypertensive mouse model. If this is the case, the contraction could occur by at least two distinct pathways. Firstly, the WNK1/WNK3/SPAK/OSR1 pathway is thought to be important for regulating vascular tone by controlling the phosphorylation state of the NKCC1 co-transporter and hence the membrane potential of vascular smooth muscle (VSM) cells (Yang et al., 2010; Bergaya et al., 2011; Zeniya et al., 2013). So if either WNK1 or WNK3 accumulates in the vessel wall of the CUL3WT/A403–459 mice, the contractile tone would be expected to rise through depolarisation of VSM and increased calcium entry. Alternatively, the mutant Cullin-3 protein may directly affect the phosphorylation state of myosin light chain by regulating RhoA/RhoA kinase (ROCK) levels (Ibeawuchu et al., 2015). RhoA protein levels in the VSM are thought to be regulated by CUL3 in complex with the substrate adaptor RhobBT1 (Pelham et al., 2012). If these pathways are operational in PHA2E, then they suggest that treatment with a thiazide diuretic or a specific ROCK inhibitor (Liao et al., 2007) may be the more appropriate drug of choice. Central aortic pressure elevation is now widely accepted to be an important and independent cardiovascular risk factor (Liao & Farmer, 2014), so if it is elevated in patients with PHA2E, it will be important to show that it is normalised by appropriate pharmacotherapy.

Materials and Methods

All plasmids, antibodies and recombinant proteins that we have generated for this study are available on request from the MRC-PPU reagents website (http://mrcppureagents.dundee.ac.uk/).

Plasmids and protein purification

The following plasmids and/or protein purifications have been described elsewhere in Ohta et al. (2013): KLHL3 (DU23218), DACTEV-CUL3-RBX1 (DU23291), His-UBE1 (DU32888) and His-TEV-UBE2D3 (DU15703); and in Kelsall et al. (2013): Nedd8 (DU20689), UBE2M (DU15804) and APPBP1/UBA3 heterodimer (Nedd8 activating E1) (DU21784).

DACTEV-CUL3/A403–459_RBX1 (DU23292) was cloned in a comparable way to DACTEV-CUL3WT_RBX1 (DU23291). Specifically, the pFastBac Dual DACTEV expression system was created by subcloning a BglII-BamHI-flanked PCR product encoding the full-length DACTEV tag followed by a TEV protease site into a

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Figure 7.

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Cullin-3 and defective WNK ubiquitylation in PHA2E  
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<table>
<thead>
<tr>
<th>Parameter</th>
<th>CUL3&lt;sub&gt;WT&lt;/sub&gt; (n = 10)</th>
<th>CUL3&lt;sub&gt;WT/A403-459&lt;/sub&gt; (n = 11)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wall thickness (µm)</td>
<td>99.6 (± 3.5)</td>
<td>120.8 (± 3.3) ***</td>
</tr>
<tr>
<td>No. of elastic laminae</td>
<td>5.3 (± 0.1)</td>
<td>5.4 (± 0.1)</td>
</tr>
</tbody>
</table>
pFastBac™ Dual (Life Technologies, UK). Human Cullin-3 (A403–459; GenBank NM_003590.4) was codon-optimised for expression in insect cells and custom synthesised (GenScript USA Inc.) before being subcloned downstream of the Dac-tag in cassette two of this vector. Human RBX1 (GenBank NM_014248.2) was amplified from an EST IMAGE clone 3138751 and subcloned into pGEX6P-2 (GE Healthcare LifeSciences UK) for a Not1 flanked ORF from EST IMAGE clone 5265409 and an EST IMAGE clone 5265409 and subcloned into cassette one (unvector. Human RBX1 (GenBank NM_003590.4) was codon-optimised for expression in GST-tagged fusion protein in BL21 cells. Expression was induced 150 mM NaCl and 10% glycerol.

The fusion protein was then concentrated and further purified by size-phase by incubation with C-terminally His-tagged TEV protease sepharose. After washing, the protein was recovered from the solid was captured from the lysate by incubation with ampicillin-Insect-Protein-free Insect Cell Medium (Lonza). The fusion protein CUL3 and untagged RBX1 (Lee et al., 2012) was expressed as an N-terminally His antibody were purchased from Sigma-Aldrich. The rabbit anti-ubiquitin (AB7254) were purchased from Abcam. The goat anti-GFP (AB5450) and the mouse monoclonal anti-SpaK phospho-Thr44 antibody were purchased from Biosource. The rabbit anti-phospho-Thr44 antibody (residues 38–76 of mouse SPAK, S668D), SPAK-mouse C-terminal antibody (424–556 of mouse SPAK, S669D), SPAK/OSR1 (T-loop) phospho-Thr233/Thr185 antibody (226–238 of human SPAK or residues 178–190 of human OSR1, TRNKVRFpTFVGTP, S204C), SPAK-phospho-Thr233 antibody (226–238 of human SPAK, TRNKVRFpTFVGTP, S668B and S668D), NKCCI-total antibody (residues 1–260 of shark NKCCI1, S841B), NCC-phospho-Thr60 antibody (residues 54–66 of human NCC phosphorylated at Thr60, RTFGYNpTDIVVpPT, S959B), NCC-phospho-Thr44 antibody (residues 38–52 of mouse NCC phosphorylated at Thr44, SQPSHTpHGSTLYMRM, S242C) and KLHL3 N-terminal mouse (1–21 of mouse KLHL3, S740D). The anti-ERK1/2 antibody (9102) was purchased from Cell Signaling Technology. The anti-FLAG antibody (F1804), anti-β-actin antibody and anti-His antibody were purchased from Sigma-Aldrich. The rabbit anti-human CUL3 antibody was a kind gift from Izabela Sumara at the IGBMC, Strasbourg (Sumara et al., 2007). The following commercial rabbit anti-human antibodies were obtained and used for immunodetection: KLHL3 antibody (HPA01291) from Atlas Antibodies; KLHL3 (AB196776) (~14% immunogen peptide homology with KLHL2), NCC-total [SLC12A3] (AB95302) and LAMPI (AB24170) from Abcam; and PV25 (PVALB) from Swant.

Antibodies

The following antibodies were raised in sheep and affinity-purified on the appropriate antigen by the Division of Signal Transduction Therapy Unit at the University of Dundee: WNK1-total antibody (residues 2,360–2,382 of human WNK1, S62B), WNK4 N-terminal antibody (residues 1–14 of mouse WNK4, S726D), SPAK-mouse N-terminal antibody (2–76 of mouse SPAK, S668D), SPAK-mouse C-terminal antibody (424–556 of mouse SPAK, S669D), SPAK/OSR1 (T-loop) phospho-Thr233/Thr185 antibody (226–238 of human SPAK or residues 178–190 of human OSR1, TRNKVRFpTFVGTP, S204C), SPAK-phospho-Thr233 antibody (226–238 of human SPAK, TRNKVRFpTFVGTP, S668B and S668D), NKCCI-total antibody (residues 1–260 of shark NKCCI1, S841B), NCC-phospho-Thr60 antibody (residues 54–66 of human NCC phosphorylated at Thr60, RTFGYNpTDIVVpPT, S959B), NCC-phospho-Thr44 antibody (residues 38–52 of mouse NCC phosphorylated at Thr44, SQPSHTpHGSTLYMRM, S242C) and KLHL3 N-terminal mouse (1–21 of mouse KLHL3, S740D). The anti-ERK1/2 antibody (9102) was purchased from Cell Signaling Technology. The anti-FLAG antibody (F1804), anti-β-actin antibody and anti-His antibody were purchased from Sigma-Aldrich. The rabbit anti-human CUL3 antibody was a kind gift from Izabela Sumara at the IGBMC, Strasbourg (Sumara et al., 2007). The following commercial rabbit anti-human antibodies were obtained and used for immunodetection: KLHL3 antibody (HPA01291) from Atlas Antibodies; KLHL3 (AB196776) (~14% immunogen peptide homology with KLHL2), NCC-total [SLC12A3] (AB95302) and LAMPI (AB24170) from Abcam; and PV25 (PVALB) from Swant. The goat anti-GFP (AB5450) and the mouse monoclonal anti-ubiquitin (AB7254) were purchased from Abcam. The MYPT1 phospho-Thr696 mouse monoclonal antibody (MAB0001) was obtained from Abnova. Secondary antibodies coupled to horseradish peroxidase used for immunoblotting were obtained from

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Pierce. Secondary antibodies conjugated to fluorochromes for LiCor Odyssey Western blot scanning were obtained from Licor (www.licor.com) and Life Technologies. Fluorochrome-conjugated secondary antibodies for immunofluorescent confocal microscopy were obtained from Life Technologies and Abcam.

In vitro assays

Ubiquitin E2 discharge assays were performed based on the method described (Plechanovová et al., 2011). Firstly, His-UBE2D2 (150 μM) was incubated in the presence of 2 mM ATP, 5 mM MgCl₂, 200 μM ubiquitin and 0.25 μM UBE1 in 50 mM HEPES pH 7.5, 150 mM NaCl, 10% glycerol and 1 mM DTT at 16°C for 15 min. To stop the reaction, we depleted ATP by incubating the charging mixture with apyrase (4.5 U/ml; New England Biolabs) at 16°C for 10 min. To this, 1 μM E3 was added and discharge reactions were then allowed to proceed for the time indicated in the figure, at 30°C. Reactions were stopped by the addition of non-reducing SDS-Laemmli sample buffer. 16% bis-acrylamide non-reducing SDS-PAGE, followed by immunodetection with anti-His antibody, allowed visualisation of ubiquitin discharge. E3 ubiquitin-ligase activity assays were based on those described previously in Ohta et al. (2013). When used as a substrate in assays, full-length WNK proteins were purified from HEK-293 cells, and either 5 μl of immunoprecipitated endogenous WNK1 protein derived from 0.5 mg of HEK-293 cells or 5 μl of immunoprecipitated over-expressed FLAG-WNK4 from 0.05 mg of HEK-293 cells was used for each assay reaction. Ubiquitylation assays typically contained 20 mM HEPES/HCl (pH 7.5), 150 mM NaCl, 2 mM DTT, 10% (w/v) glycerol, 8 μM CUL3-RBX1 complex (WT or mutant), 7 μM KLHL3, 7 μM UBE1, 60 μM UBE2D3 and 3,000 μM ubiquitin. The concentrations of some components were altered on occasion (e.g. Fig 3B) to best illustrate a given point. Reactions were initiated by adding ATP and MgCl₂ to a final concentration of 1 mM, and samples were incubated for the times indicated at 30°C. Reactions were stopped by the addition of SDS sample buffer, and samples were analysed after SDS-PAGE followed by immunoblotting or Coomassie blue gel staining. For the N8-ligase activity assays, 1 μM E3 was incubated at 30°C with 68 μM Nedd8, 8 μM UBE2M and 0.2 μM NAE in the presence of 0.15 mM ATP, 1.5 mM MgCl₂, 50 mM HEPES pH 7.5, 150 mM NaCl and 20% (w/v) glycerol for the time indicated. Reactions were stopped by the addition of SDS sample buffer, and samples were analysed following SDS-PAGE separation and Coomassie blue staining.

In vitro protein interaction pull downs

For KLHL3-CUL3, anti-KLHL3 resin was made by covalently coupling anti-KLHL3 antibody (S377D) to Protein G sepharose utilising dimethyl pimelimidate. Anti-KLHL3 resin, or control resin (prepared in the absence of any antibody), was incubated with saturating amounts of purified KLHL3 and washed thrice, resulting in KLHL3 resin. This resin was then incubated with CUL3WT or CUL34403-459 at 4°C for 1 h, before being washed thrice with PBS 0.02% Tween-20. For CAND1-CUL3, purified GST-CAND1 or GST was incubated in excess with resin at 4°C resulting in GSH-resin-GST or GSH-GST-CAND1 resin. Each resin was then incubated with purified CUL3WT or CUL34403-459 for 1 h at 4°C before being washed once with PBS 0.02% Tween-20 and then twice with PBS in the absence of detergent. Following washing, volumes were reduced to a minimum and proteins were eluted from the resin by the addition of SDS sample buffer, prior to analysis by SDS-PAGE and immunoblotting.

Cell culture

HEK (human embryonic kidney)-293 cells were cultured in 14-cm dishes in DMEM (Dulbecco’s modified Eagle’s medium; Life Technologies) supplemented with 10% (v/v) foetal bovine serum, 2 mM l-glutamine, 100 units/ml penicillin and 0.1 mg/ml streptomycin. Protein expression was induced for 24 h with 1 μg/ml tetracycline (Life Technologies). To obtain endogenous WNK1 for activity assays, we followed the immunoprecipitation protocol described in Ohta et al. (2013). To obtain wild-type FLAG-WNK4, we utilised the cell lines from Schumacher et al. (2014) and Ohta et al. (2013). When FLAG-WNK4 was to be used for in vitro ubiquitylation assays, a 14-cm plate of confluent cells was lysed in 0.5 ml ice-cold mammalian lysis buffer A (mammalian lysis buffer A: 50 mM Tris/HCl (pH 7.5), 0.15 M NaCl, 1 mM EGTA, 1 mM EDTA, 1 mM Na₂VO₄, 50 mM NaF, 5 mM Na₃P₂O₇, 0.27 M sucrose, 1% (w/v) Nonidet P-40, 1 mM benzamidine, 0.1 mM PMSF, 0.1% 2-mercaptoethanol, and Roche protease inhibitor mix (1 tablet in 50 ml) for M2 affinity purification). Lysates were clarified by centrifugation, and to this, 10 μl M2 resin per 100 μl lysate was added. Incubation of the resin with the cellular lysate for 1 h at 4°C was followed by two washes with mammalian lysis buffer A, and two further washes with 1× PBS to give a pure M2-Flag-WNK4 slurry. For interaction studies with FLAG-CUL3WT and FLAG-CUL34403-459, the same method was followed as above, with an alternative mammalian lysis buffer B (50 mM HEPES/KOH pH 7.2, 5 mM Mg(C₂H₃O₂)₂, 70 mM KC₂H₃O₂, 0.2% (w/v) Triton X-100, 10% (w/v) glycerol, 0.2 mM EDTA, 1 mM Na₂VO₄, 50 mM NaF, 5 mM Na₃P₂O₇ and Roche protease inhibitor mix (1 tablet in 50 ml)). When it was desirable to prevent deneddylation, buffers were supplemented with 50 μM 1,10-phenanthroline.

Mass spectrometry analysis

Prior to SDS-PAGE, samples were reduced and alkylated in the following way. Reactions were stopped at the time shown and incubated at 95°C in 1× LDS (Invitrogen) 5 mM DTT for 1 min. Alkylation occurred at room temperature for 30 min by the addition of 20 mM iodoacetamide (20 mM) to the samples. To quench the reaction, DTT was added to a final concentration of 20 mM, and the sample was then processed on precast 4–12% gradient gel (Invitrogen). Gel pieces were excised as shown by the boxed areas in the figure and in gel digestion of the proteins with 5 μg/ml trypsin and subsequent analysis. Mass spectrometric analysis was performed by LC-MS-MS on a linear ion trap–orbitrap hybrid mass spectrometer (Orbitrap-VelosPro; Thermo) coupled to a U3000 RSLC HPLC (Thermo). Peptides were trapped on a NanoViper trap column, 2 cm × 100 mm C18 5 mm 100 Å (Thermo, 164564) and then separated on a 15-cm Thermo EasySpray column (ES800) equilibrated with a flow of 300 nL/min of 3% solvent B [solvent A: 2% acetonitrile, 0.1% formic acid and 3% DMSO in H₂O; solvent B: 80% acetonitrile, 0.08% formic acid and 3% DMSO in H₂O]. The elution
gradient was as follows: time (min): solvent B (%): 0:3, 5:3, 45:35, 47:99, 52:99, 55:3 and 60:3. The instrument was operated with the “lock mass” option to improve the mass accuracy of precursor ions, and data were acquired in the data-dependent mode, automatically switching between MS and MS-MS acquisition. Full-scan spectra (m/z 400–1,600) were acquired in the orbitrap with resolution R = 60,000 at m/z 400 (after accumulation to an FTMS Full AGC Target, 1,000,000; FTMS Msn AGC Target, 50,000). The 20 most intense ions, above a specified minimum signal threshold (2,000), based upon a low-resolution (R = 15,000) preview of the survey scan, were fragmented by collision-induced dissociation and recorded in the linear ion trap (Full AGC Target, 30,000; Msn AGC Target, 5,000). Data files were analysed by Proteome Discoverer 1.4-SP1 (Thermo), using Mascot 2.4.1 (www.matrixscience.com), and searching an in-house database containing the relevant sequences. Scaffold (www.ProteomeSoftware.com) was also used to examine the Mascot result files. Allowance was made for the following fixed, Carbamidomethyl (C), and variable modifications, Oxidation (M), Dioxidation (M), GlyGly (K), LeuArgGlyGly (K) and Phospho (ST). Error tolerances were 10 ppm for MS1 and 0.6 Da for MS2.

Generation and genotyping of WT and CUL3

Standard housing conditions were used to maintain the mice. Taconic Artemis generated the CUL3

mice on a C57BL/6N background, which were backcrossed onto C57BL/6J to maintain the colony. All experiments were performed using littermates that were backcrossed at least once onto C57BL/6J. Genomic DNA from ear biopsies was isolated and used to genotype all mice in the study. Genotyping by PCR utilising primer 1: AAACTTACCCACTTGTTTGCC, and primer 2: AGACATCTCAGGTATCTATGGGC detected the presence of the CUL3WT (683 bp) or CUL3

allele. Studies were conducted on groups of approximately equal ratios of male-to-female mice with the exception of in vivo cardiovascular phenotyping in which a 1:2 ratio of male-to-female mice was used. All animal studies and breeding were approved by the University of Dundee ethical committee and performed under a UK Home Office-approved licence.

Mouse kidney and testes lysates

Mouse tissues were rapidly harvested and either snap-frozen in liquid nitrogen and stored at −80°C or incubated in 1 ml RNalater® solution (Sigma-Aldrich) at 4°C for 16–30 h before being removed from this buffer, patted dry with clean tissue and then stored dry at −80°C. Lysates were prepared by homogenising tissue on ice using a Polytron PT1200C homogenizer (Kinematica) in 2 ml ice-cold mammalian lysis buffer and then clarified by centrifugation at 18,000 g for 30 min. The clarified supernatant was then quantified and used immediately. Any excess sample was snap-frozen in liquid nitrogen in single-use aliquots for use as required.

The lysates were used for immunoprecipitation. Immunoprecipitations were prepared in the same way, with the following exceptions, all designed to promote deneddylation of the cullin proteins. Detergent was not included in the lysis buffer. Following homogenisation, samples were incubated on ice for 4 h. Following clarification by centrifugation at 18,000 g for 30 min, samples were aliquoted into fractions and either frozen in liquid nitrogen and stored at −80°C or used immediately. Clarified supernatant was incubated at room temperature for 1 h to enable further deneddylation, and samples were then cooled on ice and centrifuged a second time to ensure any precipitate was removed. Anti-CUL3 antibody (S067D) cross-linked (DMP) to Protein G agarose was mixed with saturating amount of lysate and incubated rotating overnight at 4°C. The next morning, the agrose beads were washed thrice with mammalian lysis buffer B and twice with 1× PBS. Proteins were eluted from beads by SDS sample buffer addition and boiling at 95°C for 10 min prior to centrifugation and SDS–PAGE and immunodetection.

Immunoblotting

Lysates or protein mixes were boiled at 95°C with 1× SDS–Laemmli sample buffer, for 5 min before being subject to SDS–PAGE (8 or 10% Tris-glycine gel, or 4–10% gradient gels (Life Technologies) or self-prepared 16% bis-acrylamide). Gels were transferred to nitrocellulose membrane using the standard wet transfer method (15% MeOH Towbin transfer buffer). Following blocking of the membranes (TBST 5% (w/v) dried skimmed milk), they were then incubated overnight at 4°C with the primary antibody indicated. All DTTT-produced sheep antibodies were used at 2 μg/ml, and phospho-specific antibodies included the addition of 10 μg/ml of the dephospho-peptide used to raise the antibody. Commercial antibodies were diluted 1 in 1,000. Membranes were then washed in TBST five times and incubated with the appropriate secondary HRP antibody at room temperature at 1:5,000 dilution. Membranes were subsequently washed five times, and the HRP signal was detected using chemiluminescence reagent (Pierce). Immunoblots were developed using an automatic film processor (SRX-101; Konica Minolta Medical).

CUL3 structural docking

Cullin-3 was modelled using Phyre2 server (Kelley & Sternberg, 2009) using human CUL3

as the sequence input and Cullin-1 chosen for the docking model (Q13616, PDB:1LDK; Zheng et al, 2002). A coil was predicted in place of the missing structural elements. UBE2D–UB was docked to RBX1 using PDB: 4AP4. Images were modified and made in UCSF Chimera (Pettersen et al, 2004; http://www.cgl.ucsf.edu/chimera).

Immunoprecipitation (mouse tissue and cell extracts)

Lysates of cellular preparations (buffer B) from FLAG-tagged CUL3 were mixed with M2 resin in the presence of 50 μM 1,10-phenanthroline to prevent deneddylation and were incubated rotating at 4°C for 1 h. Resins were washed twice with mammalian buffer B and twice with 1× PBS, the volume was then reduced to ~15 μl, and following the addition of SDS gel sample buffer to 1×, samples were boiled and the entire sample was loaded to allow analysis by SDS–PAGE and immunodetection with the antibodies indicated.

Human and mouse aorta sample preparation

Human aortic tissues were collected from donors through transplant coordinators at Addenbrooke’s Hospital, Cambridge. All samples were handled in accordance with the policies and
were then loaded onto a Bolt/C226 (Novex×#BG04125BOX) and run at 165 V in 1× MES running buffer (Life Technologies #78833) lysis buffers containing protease and phosphatase inhibitors (Roche #1183617001 and #04906845001). All steps were carried out at 4°C. Protein concentrations were determined with the Pierce™ BCA protein assay (Life Technologies #23225).

Human and mouse aorta immunoblotting

A total of 10 µg of aortic protein lysates was separated by SDS-PAGE. Prior to loading, samples were heated at 70°C for 10 min in 1× LDS sample buffer and 1× reducing agent (Life Technologies #B0007 and # B0009) to a total volume of 20 µl/well. Samples and 5 µl of Precision Plus molecular weight ladder (Biorad #161-0374) were then loaded onto a Bolt® 4–12% Bis–Tris Plus Gel (Novex® #BG04125BOX) and run at 165 V in 1× MES running buffer (Novex® #B0001), for 35 min or until the dye front reached the end of the gel. Resolved proteins were then transferred to 0.22 µM nitrocellulose membrane (Life Technologies #IB23001) using the iblot2 (Life Technologies) at 20 V for 7 min. Antibodies were incubated in 5% milk. Membranes were incubated in 5% milk or BSA in TBS for 1 h at room temperature. Primary antibodies were incubated overnight at 4°C in 5% milk or BSA in TBS–TWEEN (0.1% v/v Tween) and then washed 6× in TBS–TWEEN. Secondary antibodies were incubated in TBS–TWEEN for 1 h at room temperature in the dark and then washed 6× in TBS–TWEEN. Membranes were imaged and integrated intensity values quantified (bands were normalised against β-actin) using the LiCor Odyssey system (www.licor.com).

Sample preparation and immunostaining for imaging

Formal-fixed, paraffin-embedded human tissue sections were obtained from the Cambridge Human Research Tissue Bank. Harvested mouse tissues were immersion-fixed in fresh 4% w/v formaldehyde–PBS pH 6.9 for 16 h at 37°C and washed 3× in PBS and stored at 4°C until paraffin embedding. Five-micrometre sections were deparaffinised in Histoclear (National Diagnostics) and rehydrated in graded methanol steps. An antigen retrieval step was performed with R-Universal buffer in the 2000 antigen retriever for a single heat-pressure cycle (Aptum Biologics). Sections were permeabilised with 0.05% v/v Triton X-100–PBS for 20 min and blocked for 1 h at 37°C with 2% v/v donkey serum in 0.05% v/v Triton X-100–PBS. Primary antibodies were incubated overnight in 16 h at 4°C at the following concentrations diluted in 1% v/v donkey serum in 0.05% v/v Triton X-100–PBS: 2 µg/ml for pNCC T44, INCC, CUL3 (Sumara et al., 2007); 4 µg/ml for KLHL3 N-terminal, WNK4 N-terminal, SPAK C-terminal and pSPAK T233 (to detect mouse pSPAK T243); and 1 µg/ml for LAMPI, 1:500 for ubiquitin and 1:2,000 for PVALB. Phospho-specific antibodies included the addition of 10 µg/ml of the non-phospho form of the peptide used to raise the antibody per 2 µg/ml of antibody used. Slides were then washed for 20 min in 0.05% v/v Triton X-100–PBS and incubated in secondary antibody for 1 h at 37°C. Pre-absorbed donkey IgG-conjugated Alexa Fluor 488, 568, 633 and 647 secondary antibodies (Life Technologies/Abcam) were used at 1:1,200 dilution in 1% v/v donkey serum in 0.05% v/v Triton X-100–PBS for immunofluorescence labelling. Slides were washed as described above, mounted using Prolong Gold Antifade (P36930; Life Technologies) and shielded from light. Human IHC DAB staining was performed with ImmPRESS Reagent peroxidase universal anti-mouse/rabbit IgG (MP-7500, Vector) kit, after which slides were then dehydrated through graded methanol steps and mounted with DPX.

Image acquisition and processing

Immunofluorescence images were acquired on the Leica TCS SP2 laser scanning confocal microscope with 488-, 543- and 633-nm laser lines mounted on an upright Leica DM RXA fluorescence microscope using a HCX PL APO 63×/1.40NA oil immersion objective. Acquisition parameters were as follows: 12-bit, 1,024 × 1,024 pixels, 2.6× digital zoom, 800 Hz scan speed, 4-line Kalman filtering, sequential (by line) channel imaging, and 10-slice z-stack of 5 µm. Immunohistochemically stained specimens were imaged in brightfield on an Olympus BX51 upright epifluorescence microscope using a UPLANFL 20×/0.5NA dry objective with the Infinity3 (Lumenera) CCD set to 1,936 × 1,456 pixels.

In FIJI image analysis software (http://fiji.sc/Fiji), fluorescent z-stacks underwent background subtraction (1,000-pixel-radius rolling ball, no smoothing) and maximum-intensity z-projection. Brightness and contrast were adjusted by linear histogram stretching to enhance visibility. Any images to be compared to one another were processed in parallel. Similarly, brightfield images underwent linear histogram stretching to increase visibility uniformly across the image.

Morphometric analysis

Body weight was regularly measured two times per week throughout the study to determine differences in mass in age-matched mice. Body length was measured using photographs of mice post-mortem with a reference scale. Full body length was determined as well as tail:body length ratio. Three 5-µm sections of thoracic aorta were taken at upper (heart), mid and lower (near diaphragm) segments per mouse. These were deparaffinised, rehydrated and stained with haematoxylin and eosin, and then dehydrated and mounted. Each section was imaged under brightfield illumination and epifluorescent imaging of autofluorescent elastin laminae with a GFP filter set. Using FIJI image analysis software, macros were scripted to draw eight lines at ~22.5° to one another that intersect the aortic ring at a total of 16 points. At the point of intersection on the outer circumference, the distance from the external lamina to the inner lumen was measured. The average of the three rings (max of 16 points per ring, with points excluded if they fell on an arterial branching point or incomplete portion of a ring) was used to calculate the average thoracic aorta intima-media wall thickness. The above step was repeated to count the average number of elastic laminae in the thoracic aorta of each mouse. Data collection and analysis were carried out in a blinded fashion throughout.
In vivo cardiovascular physiology

Animals were anaesthetised with isoflurane on 100% O₂ (induction: 3%, maintenance: 1.75%) and placed on a self-regulating rectal probe-coupled heat mat (TC-1000; CWE) to maintain a body temperature of 37°C. Heart rate was measured by the R-R wave interval from ECG Lead II using an animal bio amp (FE136; AD Instruments) with needle electrodes inserted into fore and hind limbs (MLA1213; AD Instruments). The right carotid artery was catheterised with a 1F Mikro-Tip pressure transducer (SPR-1000; Millar) connected to a bridge amp (FE221; AD Instruments) and powerlab system (PL3504/P; AD Instruments). When the animals had stabilised, measurements were taken at 2,000 samples/s using LabChart version 7/8 Pro (AD Instruments) to record ECG and blood pressure pulse waveforms. In vivo dose–response to vasopressor agents were taken after baseline blood pressure traces were obtained from mice. The right external jugular vein was cannulated for administration of bolus doses in increasing half-log steps of μg/kg body weight (bw); initially, phenylephrine was administered, and after washout and return to baseline, angiotensin II was administered in a similar fashion. Each dose concentration was made up as a half-log serial dilution in sterile 0.9% w/v saline and administered at a volume of 0.2 ml/kg bw with an accompanying 20 μl 0.9% w/v saline cannula flush between doses.

Data processing and analysis were performed in LabChart 8 Pro. Using the blood pressure add-on, systolic, diastolic and dicrotic notch blood pressures were automatically detected per beat (beats with respiratory-induced artefact were gated out of the analysis using the beats classifier). Mean arterial pressure was calculated as (1/3 peak systolic pressure + 2/3 end diastolic pressure) and pulse pressure as (peak systolic pressure – end diastolic pressure). A macro was scripted to detect the anacrotic notch, by using the third zero value crossing the fourth derivative of the blood pressure, as described in Kelly et al. (1989). The augmentation pressure was calculated as (peak systolic pressure – anacrotic notch pressure) and augmentation index as (augmentation pressure/pulse pressure). A macro was scripted to measure the slope of the diastolic pressure decay, 30 ms after the dicrotic notch and 20 ms before the end diastolic pressure (to avoid perturbations caused by aortic valve opening/closing). The reciprocal of this slope (ignoring the sign and calculated from the 20% trimmed mean values) was used to determine the time decay constant of the diastolic pressure decay which correlates with vascular resistance, as described previously (Bourgeois et al., 1974). Dose–response curves were generated and data analysed (GraphPad Prism 5 and LabChart 8 Pro) using a 4-parameter logistical function to determine the estimated dose producing half-maximal response (ED50) and maximum response (Emax). Data collection and analysis were carried out in a blinded fashion throughout. Sample sizes for the mouse work were decided based on previous experience in a related mouse model (Rathi et al., 2010). When groups of animals were being compared, procedures were performed on animals selected at random from the groups.

Blood and urine analyses

To reduce variability introduced by dietary potassium ingestion, animals were fasted for a minimum of 4 h before being anaesthetised with isoflurane on 100% O₂ (induction: 3%, maintenance: 1.75%) and placed on a self-regulating rectal probe-coupled heat mat (TC-1000; CWE) to maintain a body temperature of 37°C. To minimise air contact of arterial bloods during measurements, the right carotid artery was cannulated with a mouse arterial catheter (MAC-02; SAI Infusion Technologies) pre-flushed with 10 U heparin-0.9% w/v saline to prevent clots. Blood was released via the catheter into the EC8+ cartridge for analysis on the iSTAT (Abaxis). Post-surgery additional blood was collected by using Microvette® CB 300 LH (Sarstedt) and centrifuged at 2,000 g for 5 min to extract plasma before storage at −80°C. Plasma aldosterone was calculated by HTRF (homogeneous time-resolved fluorescence) aldosterone assay (Cisbio, Codeolet, France), according to the manufacturer’s protocol using the PheraStar FS (BMG Labtech) plate reader.

To determine plasma and urine electrolyte homoeostasis, animals were placed on a 0.3% w/w Na diet for 2 weeks with urine and plasma time-matched samples collected between days 7 and 10 for normal Na (NNa) diet baseline. On day 14, mice were switched onto a 0.03% w/w Na diet with sampling repeated on days 7–10 for low-Na (LNa) diet measurements. Spot urine was collected from awake mice following spontaneous micturition on handling over a sheet of Saran® wrap or Parafilm®. Samples were then divided, and one sample received acidification with HNO₃ to a final concentration of 1% v/v to prevent precipitation of electrolytes, before both were stored at −80°C. Blood was collected by saphenous venepuncture in awake restrained animals, and plasma was separated and stored in a similar fashion to that described above. Plasma and urine (non-acidified) creatinine levels were assayed in the Core Biochemical Assay Laboratory, Addenbrooke’s Hospital, Cambridge, UK. Plasma and urine (acidified) samples were diluted 1:1,000 using ultrapure polished water containing 1% v/v HNO₃. Cations were then measured by inductively coupled plasma-optical emission spectrometry (Perkin Elmer ICP-OES Analyser) with known concentration standards and preset elemental spectra. Data collection and analysis were carried out in a blinded fashion throughout.

Statistical analysis

Data presented are mean ± standard error of the mean (SEM), unless otherwise stated. Statistical analyses were performed on either SPSS version 22 or GraphPad Prism® (http://www.graphpad.com/scientific-software/prism/) software. Two-tailed Student’s t-tests were performed either paired or unpaired where appropriate. P < 0.05 was taken as statistically significant throughout. For MYPT1 phosphorylation quantification (Appendix Fig S6), ratiometric expression was calculated for CUL3WT/A03–459 versus CUL3WT on each Western blot of independent biological replicates. The mean of the ratios and bounds of the 95% confidence interval were calculated, where ratio = 1 represents no change in phosphorylation and ratio > 1 represents an increase in the phosphorylation status of CUL3WT/A03–459 mice. Statistical significance was determined by the ratio t-test (http://www.wormbook.org/chapters/www_statisticalanalysis/statisticalanalysis.html).

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The paper explained

Problem
Mutations affecting CUL3 have been identified in patients with a syndrome of familial hypertension and hyperkalaemia. CUL3 forms a RING E3 ubiquitin ligase that controls the degradation of WNK kinases, which regulate salt homeostasis. We wanted to understand whether and how the disease-linked CUL3 mutations (that delete residues encoded by exon 9 from the protein) affect the function of the protein and the signalling through the WNK kinase cascade.

Results
The disease-linked form of CUL3 fails to ubiquitylate WNK kinase, but retains the ability to transfer ubiquitin from a charged E2, leading to ectopic self-modification of the ligase complex. Our experiments suggest that the defect in WNK ubiquitylation is likely due to increased structural flexibility of the CUL3 scaffold. The mutant form of CUL3 also fails to bind to two of its key regulators, CAND1 and the CSN. Heterozygous mice expressing the disease-linked form of CUL3 show increased signalling through the WNK kinase pathway, high blood pressure and a pattern of deranged electrolytes that is typical of human subjects with PHAE.

Impact
We provide the first detailed biochemical characterisation of a disease-causing CUL3 mutation involved in blood pressure regulation. We have also identified a previously unrecognised vascular phenotype in the mice expressing the mutant form of CUL3, suggesting they have increased vascular stiffness. If a similar phenotype exists in human subjects with the mutation, it may explain the severity of the PHAE genotype. As vascular stiffness is an important and independent risk factor for cardiovascular events, it will be important to establish whether thiazides or other therapeutic interventions are able to reduce vascular stiffness in PHAE subjects.

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Author contributions
FRS performed all biochemical analysis and experiments shown in Figs 1–3 and 4A. KS performed immunostaining and imaging in Figs 5A and 7C. KS performed surgeries, data collection and analysis of blood pressure, pressor dose–responses and pulse waveform measurements in Fig 6 with surgical support from FRS and JZ. Post-surgery KS, FRS and JZ harvested mouse tissues. KS performed dissection of aorta and kidney samples with specific experimental requirements. FRS prepared all mouse tissues and biological samples for storage as appropriate. KS performed vascular cannulations and arterial blood biochemistry data was collected with assistance from FRS; KS performed data analysis for Fig 4D. Cj and AK purified proteins used in biochemical analysis with help from FRS. RE performed structural docking and modelling shown and RE and FRS made the figures relating to this. NW performed cloning. SEC and Y collected and processed human aortic samples. RSAM and IH performed immunoblotting in Figs 4B and 7A and B with input from KS. IH performed the aldosterone assay in Fig 4C. IH and KS analysed the data. JTF performed aortic morphometric measurements in Fig 7E. NLF performed histological preparations and staining in Figs 5B and 7D and E. KS scripted macros for data extraction and processing for Figs 5A, 6B and 7C and E. KS analysed the data and generated the figures in Figs 5B and 7D and E. FRS managed the mouse study with assistance from JZ and KS. TK and KMO conceived the study and supervised the work. FRS, TK and KMO wrote the manuscript with input from KS.

For more information
All of the constructs generated in this study, and many of the purified proteins and antibodies can be purchased at the following link: https://mrccpureagents.dundee.ac.uk
MRC-PPU Kurz webpage: http://www.ppu.mrc.ac.uk/research/?pid=1004
EMIT O’Shaughnessy webpage: http://emit.medschl.cam.ac.uk/
For more information regarding the human tissue utilised in this study please go to http://www.cuh.org.uk/tissue-bank/about
OMIN disease database: http://www.omim.org/entry/145260
Protein structures mentioned in this study can be found under the noted PDB identifier online at http://www.rcsb.org

Conflict of interest
The authors declare that they have no conflict of interest.

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