Multi-omics analyses of MEN1 missense mutations identify disruption of menin–MLL and menin–JunD interactions as critical requirements for molecular pathogenicity

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Abstract

Background: Loss-of-function mutations of the multiple endocrine neoplasia type 1 (MEN1) gene are causal to the MEN1 tumor syndrome, but they are also commonly found in sporadic pancreatic neuroendocrine tumors and other types of cancers. The MEN1 gene product, menin, is involved in transcriptional and chromatin regulation, most prominently as an integral component of KMT2A/MLL1 and KMT2B/MLL2 containing COMPASS-like histone H3K4 methyltransferase complexes. In a mutually exclusive fashion, menin also interacts with the JunD subunit of the AP-1 and ATF/CREB transcription factors.

Results: Here, we applied and in silico screening approach for 253 disease-related MEN1 missense mutations in order to select a set of nine menin mutations in surface-exposed residues. The protein interactomes of these mutants were assessed by quantitative mass spectrometry, which indicated that seven of the nine mutants disrupt interactions with both MLL1/MLL2 and JunD complexes. Interestingly, we identified three missense mutations, R52G, E255K and E359K, which predominantly reduce the MLL1 and MLL2 interactions when compared with JunD. This observation was supported by a pronounced loss of binding of the R52G, E255K and E359K mutant proteins at unique MLL1 genomic binding sites with less effect on unique JunD sites.

Conclusions: Our results underline the effects of MEN1 gene mutations in both familial and sporadic tumors of endocrine origin on the interactions of menin with the MLL1 and MLL2 histone H3K4 methyltransferase complexes and with JunD-containing transcription factors. Menin binding pocket mutants R52G, E255K and E359K have differential effects on MLL1/MLL2 and JunD interactions, which translate into differential genomic binding patterns. Our findings encourage future studies addressing the pathophysiological relevance of the separate MLL1/MLL2- and JunD-dependent functions of menin mutants in MEN1 disease model systems.

Keywords: Tumor suppressor, MEN1 syndrome, Histone H3K4 methylation, AP-1 transcription factors, ATF transcription factors, Quantitative mass spectrometry, CUT&RUN genome profiling

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Mutations of the multiple endocrine neoplasia type 1 (MEN1) tumor suppressor gene are found in patients with the MEN1 syndrome [1, 2]. MEN1 is characterized...
by the occurrence of parathyroid, pituitary and pancreatic neuroendocrine tumors (PanNETs). Sporadic PanNETs frequently harbor inactivating somatic MEN1 mutations [3]. In addition, in recent years exome and whole genome sequencing studies have revealed MEN1 gene mutations in many types of cancer, such as adrenocortical, uterine, breast and other cancers [4]. The MEN1 gene acts as a classic tumor suppressor gene in endocrine tissues: loss of function results in tumorigenesis. In other tissues, such as the hematopoietic system, the MEN1 gene has pro-oncogenic activity and protein–protein interactions of the MEN1 product, menin, are emerging as therapeutic targets (see below).

Menin is involved in transcriptional regulation as an intermediary protein linking transcription factors to co-activator and co-repressing proteins [5]. Most notably, menin has been reported as an integral component of mixed-lineage leukemia MLL1 and MLL2 (official names: KMT2A and KMT2B) containing COMPASS-like histone H3K4 methyltransferase complexes [6]. Loss of Men1-dependent histone H3K4 trimethylation contributes to Men1-related pNET development in mice [7]. Menin was co-crystallized with MLL1, the AP-1/ATF transcription factor member JunD or a MLL1–PSIP1 (PC4 and SFRS1 interacting protein 1, also known as LEDGF, lens epithelium-derived growth factor) heterodimer [8]. The interaction of menin with full-length JunD was first reported in yeast two-hybrid and in vitro pull-down experiments [9]. Menin contains a deep pocket that can bind short peptides of MLL1 or of JunD in a similar and mutually exclusive manner [8]. We performed affinity purification of menin-containing protein complexes followed by quantitative mass spectrometry, which confirmed that menin is present both in MLL1 and MLL2 containing COMPASS-like complexes and in complexes of JunD and related proteins [10].

The observation that menin is a critical cofactor for the subset of leukemias driven by chromosomal translocations involving MLL1/KMT2A motivated the development of menin–MLL inhibitors, which all target the MLL1-binding pocket of menin [11–13]. Several menin–MLL inhibitors display potent anti-leukemic activities in preclinical mouse models for MLL-rearranged and NPM1 mutant acute leukemias [14–17]. At present, several clinical trials investigate menin–MLL inhibitors in relapsed acute leukemias with promising early results [18].

The paradoxical roles of menin as a tumor suppressor for endocrine tissues and a pro-oncogenic factor for MLLr- and NPM1c-driven acute leukemias encourage a better understanding of menin function. Whereas menin inhibition provides a therapeutic perspective for certain hematopoietic malignancies, a rescue of menin function could be of therapeutic value for patients with endocrine tumors. At present, there are many unknowns with regard to loss of MEN1 function and the subsequent events leading to endocrine tumorigenesis. Whereas it has been demonstrated that the transcriptional function of menin is critical to prevent endocrine tumorigenesis, the exact molecular mechanism, i.e. how loss of menin function results in altered gene expression leading to endocrine tumorigenesis, is currently unresolved. Most MEN1 gene mutations are considered to disrupt menin function entirely: nonsense MEN1 gene mutations are predicted to result in degradation of the MEN1 mRNA or to result in insoluble menin proteins [1, 19]. Moreover, MEN1 missense mutations have been reported to yield unstable protein products that are targeted for proteasomal degradation [20, 21].

In this study, we sought to identify disease-related functionally impaired MEN1 mutated forms of menin using in silico methods based on crystal structures of free menin and of menin-containing complexes. Such mutant forms of menin could clarify the critical pathogenic requirements for non-MEN1 conditions in terms of protein–protein interactions for loss of menin function. By using ectopic expression, we carried out quantitative proteomic studies of a selection of mutant menin proteins in HeLa cells. We identify a set of disease-associated menin mutants with differential effects on MLL1 and JunD interactions, which were validated by analyzing genome localizations of wild type (WT) and the mutant forms of menin, as well as JunD, MLL1 and histone modifications.

Results
In silico analysis of disease-associated MEN1 mutations
We selected MEN1 gene mutations from a compiled list of 253 missense mutations observed in various human cancers. These missense mutations correspond to 245 unique single amino acid changes, which were subjected to in silico modeling for structural changes using menin crystal structure data (Additional file 5: Table S1). The selection criteria included predicted protein stability and predicted disruption of menin–MLL1 and/or menin–JunD interactions. Of these 245 MEN1 missense mutations 80 were expected to affect protein stability, because the mutation did not yield any fitting rotamers in our modeling. For 47 missense mutations, the decrease in protein stability was predicted from a loss of internal hydrogen bonds. 86 of the 253 studied mutations appeared to be potentially pathogenic for unknown reasons other than losing internal bonds or directly altering protein–protein interactions with the MLL1 and JunD complexes. Because of their proximity to the mapped interaction surfaces of menin and MLL1 and JunD, respectively, six missense mutations were predicted to
potentially directly alter the menin binding to MLL1 or JunD (Fig. 1A and B).

For the R52G and E255K mutants, disruption of the interaction of menin with MLL1 when in complex with LEDGF was predicted. The R322H, E359K and E366D mutations could disrupt both the interaction of menin with MLL1 and with the MLL1–LEDGF complex, while E359K and E366D were predicted to disrupt the interaction of menin with JunD in silico. The Q141R mutation was selected as it was considered to be a stable protein and to potentially result in a gain of interaction of menin with MLL1, when in complex with LEDGF. Importantly, all these mutations do not display clashing rotamers, which is predicted to prevent protein stability issues.

Characteristics and transient ectopic expression of selected MEN1 mutations
The selected MEN1 mutations are associated with different human endocrine or cancer pathologies (Fig. 1A). The R52G mutation was first identified in a MEN1 patient who developed primary hyperparathyroidism and a PanNET [22]. Menin mutations L168P, L264P and E366D have been identified in MEN1 families [23–25]. The Q141R mutation was identified in a sporadic gastrinoma without loss of heterozygosity LOH at the wild type (WT) locus, which is common in gastrinomas [26]. Menin E255K has been found in a kindred with familial isolated hyperparathyroidism, a condition considered to be a variant phenotype of MEN1 [27]. The R322H mutation was found in a sporadic glioma (TCGA-CS-5394-01). The mutation resulting in menin E359K was found in a sporadic angiofibroma, a skin tumor type known to be associated with MEN1, without LOH at the WT locus [28, 29]. The MEN1 gene mutation resulting in menin E408Q was derived from a non-small cell lung cancer sample (TCGA-22-4607-01).

Based on this and on our in silico analysis, we selected missense menin mutants R52G, E255K, R322H, E359K, E366D and Q141R for further functional studies. In addition, we included the L168P and L264P mutations, which are predicted to be unstable, as well as the E408Q mutation from a non-endocrine tumor as a control. E408Q was predicted to be stable and have no effect the menin–MLL1 and menin–JunD interactions as it is located far from the binding pocket (Fig. 1B).

Next, we determined the stability of the selected mutant forms of menin by ectopic expression as described before [21]. The nine different MEN1 mutations,
along with a vector control, were introduced into a CMV-driven expression plasmids, which were used for transient transfection of HEK 293 T cells in order to assess their expression levels in total cell lysates as measured by immunoblotting using menin antibodies. As shown in Fig. 2A, immunoblotting of transfected cell lysates supported the predicted stability of the selected mutations compared to the pCDNA transfection control, which displays endogenous menin protein levels. As expected, the L168P and L264P menin proteins were expressed at lower levels compared with WT menin, indicating a reduced protein stability as noted before [30]. Total MEN1 mRNA levels measured by RT-qPCR analysis as well menin immunoblot protein quantitation corrected for MEN1 mRNA levels were determined to assess the effects of mRNA variations on differences at the protein level. Variability of mRNA levels did not explain the observed differences in protein expression, except as expected for L168P and L264P (Additional file 1: Figure S1A and B).

**Quantitative mass spectrometry of stably expressed menin mutants**

We proceeded to assess the menin interactome using C-terminal GFP fusions of the menin mutants by quantitative mass spectrometry after GFP-based immunoaffinity purification. For this purpose, MEN1 WT and mutant cDNA fusions were expressed in stably transfected HeLa cell lines in an inducible manner. Immunoblot analyses of total lysates with menin antibodies indicated inducible and similar expression of the GFP-fused WT and mutant menin proteins (Fig. 2B) with the exception of L168P and L264P. In contrast to the HEK293T transient transfection experiment (Fig. 2A), GFP fusions of these two mutants are expressed to higher levels than WT menin. We noted that the menin-GFP proteins are expressed to lower levels than endogenous menin (Fig. 2B). Fluorescence microscopy and immunoblot analysis confirmed the nuclear presence of the GFP-menin WT protein as well as the entire set of mutants with the exception of menin L168P and L264P, which are predominantly present in the cytoplasm (Fig. 2C, Additional file 2: Figure S2A and B).

Quantitative mass spectrometry of purified WT menin-GFP from nuclear extracts identified all MLL1/MLL2 complex members (DPY30, KMT2A, KMT2B, WDR5, ASH2L, RBBP5, HCFC1, HCFC2, and PSIP1) as interactors (indicated by red dots in left panel of Fig. 3A). Also, we observed an interaction with OGT of a lower abundance than MLL1/MLL2 subunits as indicated by the relative stoichiometry scores (right panel, Fig. 3A) and as reported before [10]. In addition, JunD and related proteins ATF7 and FOS were found as significant interactors (marked by blue dots in left panel of Fig. 3A). The AP1 and ATF complex members FOSL2, ATF2 and ATF3 did not pass our stringent significance criteria. Whereas the volcano plots provide information on the significance of an interactor, iBAQ values provide a quantitative

![Fig. 2](image-url)
measure [10]. Based of iBAQ values we determined the stoichiometries of interacting proteins relative to the WT and mutant menin-GFP baits as shown in Fig. 3A and B. GFP-purification of the unstable L168P and L264P mutants, which are mostly cytoplasmic, displayed strongly reduced MLL1/MLL2 complex and JunD-related interactions (Fig. 3B). Menin Q141R displayed strongly reduced, but significant interactions with MLL1/MLL2 or JunD and the interactions displayed by menin E408Q were very similar to WT menin (Fig. 3B). Immuno-purification of menin mutants R52G, E255K and E359K mostly reduced interactions with MLL1/MLL2 subunits with a smaller effect on JunD-related interactions (Fig. 3B). Interestingly, the E366D and R322H mutants, which were predicted to disrupt the menin–MLL interactions, retained MLL1/MLL2 complex interactions. In contrast to the in silico predictions, JunD binding was retained entirely by the E366D mutation and at least partially in menin R322H and E359K complexes (Fig. 3B). Relative MLL1 and JunD binding of all menin mutants compared with WT menin is summarized in Fig. 3C. The volcano plots of the menin mutants are presented in Additional file 3: Figure S3.

Next, we compared the cytoplasmic interactions of the set of GFP-menin proteins by applying our stringent criteria (FDR = 0.01 and s0 = 2). Among the top 10 interactors of cytoplasmic WT menin are ubiquitin, the ubiquitin E3 ligase RNF126, the ubiquitin-chain binders RAD23A/B and VCP, and protein chaperones (HSPA8, HSPA1A/B, DNAJB1) (Fig. 4A). This indicates that newly synthesized WT menin-GFP fusion is subjected to ubiquitin-dependent proteasomal turnover in the cytoplasm, which is consistent with previous observations of transiently expressed WT and mutant menin proteins [21]. While in this earlier work CHIP(STUB1) was identified as the responsible ubiquitin-protein E3 ligase in HEK293T cells, our results suggest that RNF126 might be responsible for menin turnover in HeLa cells.

Subsequent analyses of the cytoplasmic interactome of the stable mutations showed similar interaction patterns (Fig. 4B and Additional file 4: Figure S4). However and as expected [30], the two predicted unstable MEN1 mutations resulting in menin L168P and L264P exhibited more pronounced HSPA8 and HSPA1A/B binding compared to the other menin mutants or to WT menin. HSPA8(HSC70) and HSPA1A/B(HSP70) are the most abundant protein chaperones in HeLa cells. These ATP-hydrolyzing chaperones act to assist in pulling nascent polypeptides out of the ribosome, proper de novo folding, assembly of protein complexes and controlling the degradation of mis-folded proteins [31]. Interestingly, BAG2 and BAG5, which act as nucleotide-exchange factors for HSPA8 and HSPA1A/B, are also among the most abundant menin (mutant) interactors. In conclusion, the increased association of HSPA8A and HSPA1A/B is consistent with mis-folding of the L168P and L284P menin mutants. However, WT menin and other mutants predicted to be stable also associate with chaperones and components of the controlling protein quality and abundance, which indicates that menin proteins are under strict surveillance for protein quality.

**Analyses of genome-wide binding of menin mutants and co-occupancy with JunD and MLL**

In order to determine the functional relevance of the interactome results, we determined genome-wide DNA binding of the GFP-tagged menin protein and the R52G, E255K and E359K mutants, that displayed a predominant disruption of the menin–MLL1/MLL2 over the menin–JunD interactions. First, we determined the genome distribution of MLL1 and of RNA polymerase II (pol II) by CUT&RUN [32], using MLL1 or RPB1-CTD antibodies, respectively, and of GFP-tagged JunD by greenCUT&RUN using a combination of enhancer-MNase and LaG16-MNase as described [33]. In both methods, we included mononucleosomal *Drosophila* DNA as a spike-in for normalization purposes [34]. The ENCODE HeLa cell ChiPseq datasets for the histone modifications H3K4me3, H3K4me1 or H3K27ac and for DNA accessibility determined by ATAC-seq were included. We found that WT menin is predominantly present at sites of active transcription as indicated by the presence of H3K4me3 and pol II as well as accessible DNA. Menin appeared to bind DNA either in a broad or a sharp pattern (Fig. 5A). The broad pattern overlaps with MLL1 and H3K4me3, whereas a number of sharp menin peaks seemed devoid of MLL1. In order to examine this further, menin peaks were classified into eight categories on the basis of its overlap with the MLL1, JunD and H3K4me3 peaks. This resulted...

![Figure 3](https://example.com)
Fig. 3 (See legend on previous page.)
Fig. 4 Quantitative proteomics of cytoplasmic GFP-tagged WT menin or mutant menin expressed in HeLa cells. **A** Volcano and stoichiometry plots for the significant interactors of WT menin-GFP from the cytoplasmic fraction. Control and GFP stand for pulldown with control agarose beads versus GFP-Trap agarose beads, respectively. Stringent criteria (FDR = 0.01, s0 = 2) were applied for significance scores. **B** iBAQ-based stoichiometries of the top 10 cytoplasmic interactors normalized to the WT or mutant menin-GFP bait protein. Results shown represent Intensity Based Absolute Quantification with whiskers indicating standard deviations. Please note that the MaxQuant search identified RPS27A and UBA52 as cytoplasmic menin inhibitors, but manual inspection of peptides assigned to RPS27A and UBA52 uniquely map to the ubiquitin part of these ubiquitin fusion proteins and no RPS27A or UBA52 specific peptides could be identified.
in eight different clusters (Fig. 5B). In the largest cluster 1 (9009 peaks, 49.9%) menin binding overlaps with MLL1, H3K4me3 and JunD. These sites are also marked by H3K27ac, pol II binding and DNA accessibility indicating that menin binding correlates with promoter activity as noted before in ChIPseq experiments [16, 17, 35]. The sharp menin peaks overlap with JunD and not always with MLL1 as indicated by cluster 3 and 4 (1776 peaks or 9.3% and 1170 peaks or 6.2%, respectively). A high percentage of the peaks in these two JunD+/−MLL1− clusters contain cognate AP1 or ATF consensuses sequences (35.5% and 49.7% with AP1 motifs, and 12.33% and 15.04% with ATF motifs within cluster 3 and 4, respectively; Fig. 5B). Cluster 2 peaks also have a high percentage of AP-1 and ATF motifs, and WT menin peaks are sharp. Although the peaks in this cluster display MLL1 co-occupancy, H3K4me3 marks were absent. In contrast, the cluster of 1 broad menin peaks overlapping with JunD, MLL1 and H3K4me3 peaks contain fewer AP1 or ATF motifs. Menin peaks, coinciding with MLL1 and H3K4me3 presence but not JunD binding were found to harbor the fewest AP-1 binding sites (clusters 5 and 6). Correlation analyses were performed for the greenCUT&RUN data of the menin mutants R52G, E255K, E359K with menin E408Q as a control. As expected, menin E408Q clustered with WT menin, while the other mutations form a different cluster (Fig. 6A). A representative coverage track is shown in Fig. 6B, which indicates that the profile of the E408Q mutant is similar to the WT menin. The other menin mutants both lost and retained binding when compared to WT menin. For global comparisons of menin–MLL or JunD interaction loss, we focused on WT menin–JunD peaks in the absence of MLL1/H3K4me3 peaks (cluster 4) and WT menin–MLL1/H3K4me3 peaks in the absence of JunD (cluster 5). We observed loss of menin R52G, E255K, E359K binding both to sites, that were co-occupied by WT menin and MLL/H3K4me3, and to sites, that were co-occupied by WT menin and JunD (Fig. 6C and D). In the JunD dominated cluster 4 the difference for menin R52G was more pronounced compared to menin E255K and E359K (Fig. 6D). Taken together, the genome localization data of the R52G, E255K, E359K menin mutants are consistent with the interactome data, and indicate that disruption of the interaction with the MLL1/MLL2- and JunD-containing complexes underwrite the molecular defects of disease-related mutations in MEN1.

Discussion

Understanding the molecular pathology caused by MEN1 gene mutations may help to elucidate the pathophysiology of MEN1-associated tumorigenesis. Our study started with an in silico screening approach based on crystal structures of menin, combined with immunoaffinity purification and quantitative mass spectrometry experiments to identify functionally impaired MEN1 gene missense mutation-derived proteins. Subsequently, we functionally validated our observations by analyzing the genome localization of selected mutant forms of menin. Using these methods, we successfully identified menin missense mutations (R52G, Q141R, L168P, E255K, L264P, R322H and E359K) that directly result in loss of the menin–MLL1/MLL2 complex and of menin–JunD interactions, indicating that disruption of this interaction is the most critical link in the molecular pathogenicity of menin protein function loss. While R52G, L168P and L264P were identified in MEN1 patients, the Q141R, E255K and E359K mutations were isolated from patients with pathologies related to MEN1 (Fig. 1A).

The R52G and E359K mutants have not been analyzed in protein interaction studies before. R52G was found in a MEN1 family of three generations [22]. The E359K mutation was found in an angiofibroma, a skin tumor type known to be affected by MEN1 gene mutations [28, 29]. The E255K mutation has been studied more extensively. This mutation was reported in a familial isolated hyperparathyroidism family and found to co-segregate with the occurrence of parathyroid adenomas in this family. Loss of the WT allele was confirmed in parathyroid adenoma tissue. E255K has been described in several studies on properties of the menin protein. It was found to have a half-life similar to WT menin supporting the observation, that it is a stable MEN1 gene mutation [21]. Apart from a previous report from our group in which we showed reduced interaction of menin E255K with the estrogen receptor alpha ligand binding domain in a yeast two-hybrid set-up, the menin E255K mutation has not been studied for its ability to interact in MLL1/MLL2 complexes or with JunD before [30].

(See figure on next page.)

**Fig. 5** Genome-wide occupancy of WT menin, MLL1, JunD, RNA polymerase II, genome-wide presence of histone marks and DNA accessibility in HeLa cells. A Normalized coverage tracks of the menin occupied regions showing broad and narrow peaks. B Normalized heatmaps indicating menin, MLL1, JunD, H3K4me3, H3K4me1, H3K27ac and pol II peaks, as well as ATACseq results categorized according to menin, MLL1 and JunD binding. The + and − signs indicate overlap and non-overlap with the peaks of WT menin, respectively. The eight categories spanning six clusters are indicated on the left and the numbers between brackets indicate the total number of menin peaks in each cluster. Pie charts showing the proportion of AP-1 and ATF motifs at menin–JunD sites in the different clusters are shown on the right side of heatmaps. Other indicates the proportion of peaks without AP-1 and ATF motifs.
Fig. 5 (See legend on previous page.)
Interestingly, within the three-dimensional structure of menin, residues R52 and E255 are in close proximity to each other (Fig. 7). Structural analyses indicate that R52, E255, and E359 are all part of the interaction surface with MLL1, and therefore, we expected loss of MLL1 interaction of these mutants. In addition to binding MLL1, E359 is also involved in the interaction with JunD. The incomplete loss of menin E359K–JunD binding (Fig. 3B) could be due to additional menin–JunD binding surfaces. For E255K and E359K we also find menin–JunD interactions are less affected compared to MLL1/MLL2 interactions. These observations are in line with the lack of clinical correlation of endocrine tumorigenesis and JunD function, for example the absence of JUND gene mutations in MEN1-related endocrine tumor types. This indicates that the loss of incorporation of mutant forms of menin into MLL1 and MLL2 histone methyltransferase complexes is a central feature of stable menin missense mutations in MEN1-related endocrine tumorigenesis.

The E366D mutation was found in a MEN1 family and localized in the MLL1 and JunD binding pocket [25]. However, E366D did not affect interactions with MLL1/MLL2 or JunD complexes, which suggests that the functional effects of this glutamate-to-aspartate conversion are rather subtle and missed in our assay system. We note that the E366D menin mutant displayed an interaction with the nuclear pore component NUP210 (Fig. 4B). NUP210 has recently been proposed to also act as a mechanosensor connecting the cytoplasmic components to the nucleus to control heterochromatin formation [36]. Possibly, loss of tumor suppressor activity by the E366D mutation relates to an acquired function linked to NUP210 binding. Similarly, R322H and E408Q isolated from a sporadic glioma and non-small cell lung cancer patient, respectively, did not affect MLL1/MLL2 or JunD complexes, which suggest that these effects are either minimal or passenger rather than driver mutations for these pathologies.
Taken together, the nuclear interactome data of our MEN1 mutant set indicate a pathological role for six out of nine menin mutations tested. In two recent studies, MEN1 gene missense mutations were analyzed in different in silico approach to predict unstable menin mutants [37, 38]. In contrast, we combined in silico predictions for stable menin mutants with transient and stable expression in HeLa cell and quantitative mass spectrometry to identify deficient functions of these mutants. Our functional analyses support the notion that restoration of menin–MLL1 interactions is a potential treatment goal in MEN1 and MEN1-related endocrine tumors [5].

The menin protein is under strict quality control mechanisms operating in the cytoplasm as indicated by its protein interactors in this cellular compartment. Eight of the top 10 interactors of WT menin either belong to the protein chaperone or the ubiquitination pathways (Fig. 4A). It is unlikely that this results from an overexpression artifact as all menin-GFP proteins are expressed to (much) lower levels than endogenous menin (Fig. 2B). It is also unlikely that interaction with the protein chaperone or ubiquitin pathways depend on the GFP moiety as similar experiments with the cytoplasmic fraction of other transcription factors do not identify these protein chaperone or the ubiquitination proteins [39]. Strikingly and as expected, the L168P and L264P instability mutants display increased interactions with components of chaperone and ubiquitination pathways. In accordance with this observation, reduced nuclear localization of a myc-tagged L264P menin mutant has been reported in the literature [40]. In contrast, the other menin mutants do not display increased ubiquitination machinery interactions, which indicates that they are similarly stable as WT menin. In the proteomic analyses, we found that the ubiquitin E3 ligase RNF126 was among the top interacting cytoplasmic proteins of wild type menin as well as menin mutants, indicating that this E3 ligase could be responsible for menin turnover in HeLa cells. This interaction has not been reported in the literature to date. Elevated expression of RNF126 has been reported as a prognostic biomarker in breast cancer, a cancer type associated with MEN1 gene mutations [35, 41]. The observed menin–RNF126 interaction motivates further research as RNF126 inhibition could stabilize menin mutant proteins to (partially) rescue functions in MEN1 mutated breast cancer.

Genome-wide DNA occupation analyses showed that menin and JunD are present together at many sites of active transcription (Fig. 5). While the menin–JunD interaction may not be critical for the molecular pathogenicity of MEN1 gene mutations, this remarkable co-existence remains intriguing and could still be relevant for menin function, besides endocrine tumorigenesis. It should be noted that cell-type-specific effects cannot be ruled out, as due to lack of a suitable endocrine cell line, we used human cervical carcinoma-derived HeLa cells for our experiments. Similar to menin, the JunD transcription factor is broadly expressed across many

Fig. 7 Projection of the selected MEN1 gene mutations menin R52G, E255K, E359K and E408Q (in green) on the tertiary structure of WT menin. Overlapping positions of the MLL1 peptide (in brown) and of the JunD peptide (in blue) are indicated.
tissues and we expect that our observations in HeLa cells are valid for many different cell and tissue types. We found that the menin binding pocket mutants R52G, E255K and E359K have differential effects on MLL1/MLL2 and JunD interactions, which translate into differential genomic binding patterns. This indicates subtle differences in MLL1/MLL2 and JunD binding, which are consistent with the structural data on MLL1 and JunD peptide binding (Fig. 7). In contrast, the E366D mutation observed in a MEN1 family was predicted to affect MLL1 and JunD binding, but it had little effect on both (Fig. 3B), which may be explained by the conservative nature of the E366D mutation.

Conclusions
Taken together, our findings underline the effects of MEN1 gene mutations in both familial and sporadic tumors of endocrine origin on the interaction of menin with MLL1 and MLL2 in COMPASS-like histone H3K4 methyltransferase complexes and with JunD-containing transcription factors. Our results encourage future studies addressing the pathophysiological relevance of the separate MLL1/MLL2- and JunD-dependent functions of menin mutants in MEN1 disease model systems. In addition to menin’s role as a tumor suppressor in MEN1-associated endocrine tissues, the MEN1 gene has prooncogenic activity in the hematopoietic system. A role for AP-1 transcription factors like JunD in MLLr and NPM1 mutant acute leukemias has not been explored yet, but the first clinical trial results with menin–MLL inhibitors combined with our menin mutant results motivate future studies of these inhibitors on the menin–JunD interaction and its possible functional involvement in acute leukemias.

Methods
Identification of disease-associated MEN1 mutations
A list of MEN1 gene missense mutations was compiled using data from the Human Gene Mutation Database (HGMD:(42)) for MEN1 mutations and the Catalogue of Somatic Mutations in Cancer (COSMIC;(43)) for somatic MEN1 mutations, complemented with data from the literature [8, 9, 20](Additional file 5: Table S1).

In silico analysis of MEN1 missense mutation-derived menin proteins
The PDB file of menin (3U84, reduced to monomer, containing 550 amino acids of the 610 amino acid menin protein) was analyzed using Swiss-PdbViewer 4.1.0. Menin protein products resulting from MEN1 mutations were assessed based on the following criteria: surface accessibility, predicted charge and polarity changes due to the mutations, presence of hydrogen bonds as well as salt bridges at the site of the mutations (using VMD 1.9.2 [44]). For every mutation, the number of rotamers, which showed clashes within the tertiary protein structure was counted (Additional file 5: Table S1). When using default software settings, the absence of clashing rotamers (using default software settings) due to the MEN1 mutations was defined as stable, less than 100% clashing rotamers as potentially stable and 100% clashing rotamers due to the mutation as predicted to result in unstable menin protein. The PDB files of the menin–MLL1 complex (3U85), menin–MLL1–LEDGF complex (3U88) and menin–JunD complex (3U86) were acquired from RCSB [45] for analysis of the effects of MEN1 mutations on the reported interactions of menin with MLL1, PSIP1/LEDGF or JunD. Mutations in the reported MLL1, PSIP1/LEDGF and JunD interaction surfaces of menin were described by selecting amino acids in a range of 5 Å from a known interaction site and verifying if any known mutations were present within this 5-Å radius. Only mutations on the surface and affecting an amino acid capable of forming H-bonds were considered for their relevance for the predicted interaction surface.

Projection of menin mutations
Both structures of the menin–JunD (ID: 3U86) and menin–MLL1–LEDGF (ID: 3U88) complexes were superimposed by sequence alignment with matchmaker and amino acids were highlighted using Chimera (version 1.13.1) [46].

Plasmids and cell lines
The MEN1 cDNA was cloned into the Gateway system using the pENTR directional TOPO cloning kit to enable C-terminal tagging (Life technologies, MA, USA). MEN1 gene mutations were introduced using site-directed mutagenesis essentially as described by the Quikchange strategy (Agilent, CA, USA) into the MEN1 cDNA in a pCDNA3.1 vector for transient expression, as well as in a pENTR-menin plasmid for C-terminal tagging with GFP via GATEWAY recombination with the pCDNA5_FRT_TO_C-GFP plasmid (van Nuland et al., 2013). The menin mutations and complete ORF of all plasmids were validated by DNA sequence analysis using M13fw and M13rev primers for the pENTR plasmids and CMV-fw and N-GFPrev primers for the pCDNA5_FRT_TO_C-GFP expression plasmids:

M13fwd: 5’-CCCAGTCACGACGTTGAAAACG-3’
M13rev: 5’-GAAAGCAGCTATGACCATG-3’
CMVfw: 5’-CGCAATATGGGCCTAGGCTGC-3’ and N-GFPrev: 5’-ACAGCTCCTCGCCCTTGC-3’.
Human embryonic kidney (HEK) 293 T and human cervical carcinoma HeLa cells were cultured in standard growth medium (DMEM, 10% FBS, 100 U/ml penicillin, 150 ug/ml streptomycin and 2 mM glutamine).

Expression of menin (mutant) proteins
HEK 293 T cells were transiently transfected with 600 ng DNA in a 12-well format using the PEI transfection reagent. After 48 h, cells were lysed in Laemml sample buffer. Total cell lysates were analyzed through immunoblotting using anti-menin (A300-105A, Bethyl) and antialpha tubulin (CP06 Calbiochem) antibodies respectively. In addition, total RNA was isolated and quantitative RT-PCR was performed to determine total MEN1 and ACTB (beta-actin) expression in pCDNA3.1 MEN1 transfected cells as described [47]. Menin protein levels were determined using ImageJ [48] from three independent transfections after subtraction of the background/menin signal from empty vector transfected cells. Ratios of relative menin protein and MEN1 mRNA levels determined in triplicate were calculated.

For doxycycline-inducible expression in stable cell lines MEN1-GFP expressing cDNAs were chromosomally integrated in HeLa cells using the Flip-in system, essentially as described [10]. After 24 h of induction using 1 µg/ml doxycycline, GFP expression was verified using immunoblotting using GFP (JL-8- Clontech) and α-tubulin (CP06 Calbiochem) antibodies. Nuclear and cytoplasmic extracts were prepared for GFP-affinity purification coupled to mass spectrometry analyses as described before [10].

Quantitative mass spectrometry of the menin interactome
GFP-affinity purification, sample preparation and data analysis were performed as reported [10, 34, 49]. Briefly, HeLa cell harboring menin-GFP cDNAs were grown in 15 15 cm dishes until subconfluency (approximately 300 million cells in total) and induced with 1 ug/mL doxycycline for 24 h. Cells were harvested by dislodging with trypsin and cell pellets washed with cold PBS (Gibco, #10,010–015). Cell pellet was re-suspended in 5 packed-cell volumes (PCVs) of ice-cold Buffer A (10 mM Hepes–KOH pH 7.9, 1.5 mM MgCl2, 10 mM KCl), incubated for 10 min on ice and then centrifuged at 400 g for 4 °C for 5 min. Supernatants was aspirated and cells were lysed in 2 PCVs Buffer A containing 1 × CPI (Roche, #11,836,145,001), 0.5 mM DTT and 0.15% NP40. The suspension was homogenized in Dounce homogenizer followed by centrifugation at 3200 g and 4 °C for 15 min. Supernatant and pellet contain cytoplasmic and nuclear fractions, respectively. The nuclear pellet was washed gently with 10 volumes of Buffer A containing 1 × CPI (Roche, #11,836,145,001), 0.5 mM DTT and 0.15% NP40 and centrifuged for 5 min at 3200 g at 4 °C min. Nuclear proteins were extracted by 2 PCVs volumes of high salt Buffer B (420 mM NaCl, 20 mM Hepes–KOH pH 7.9, 20% v/v glycerol, 2 mM MgCl2, 0.2 mM EDTA, 0.1% NP40, 1 × CPI, 0.5 mM DTT) during gentle agitation at 4 °C for 1.5 h. Both the nuclear and cytoplasmic extracts were centrifuged at 3200 g and 4 °C for 60 min. Supernatants were collected and protein concentration was measured by Bradford assay.

1 mg of nuclear or 3 mg of cytoplasmic extract was used for GFP-affinity purification as described (Spruijt et al., 2013). In short, protein lysates were incubated in binding buffer (20 mM Hepes–KOH pH 7.9, 300 mM NaCl, 20% glycerol, 2 mM MgCl2, 0.2 mM EDTA, 0.1% NP-40, 0.5 mM DTT and 1 × Roche protease inhibitor cocktail) on a rotating wheel at 4 °C in 1 h in triplicates with GFP-Trap agarose beads (#gta-200, Chromotek) or control agarose beads (#rab-20, Chromotek). The beads were washed two times with binding buffer containing 0.5% NP-40, two times with PBS containing 0.5% NP-40, and two times with PBS. On-bead digestion of bound proteins was performed overnight in elution buffer (100 mM Tris–HCl pH 7.5, 2 M urea, 10 mM DTT) with 0.1 µg/ml of trypsin at RT and eluted tryptic peptides were bound to C18 stage tips (ThermoFisher, USA) prior to mass spectrometry analysis.

Tryptic peptides were eluted from the C18 stage tips in H2O:acetonitrile (35:65) and dried. Samples were analyzed by nanoflow-LC–MS/MS with a Q-ExactivePlus mass spectrometer (Thermo Fisher Scientific) coupled to an Easy nano-LC 1000 HPLC (Thermo Fisher Scientific) in the tandem mass spectrometry mode with a 90 min total analysis time. The flow rate was 300 nl/min, buffer A was 0.1% (v/v) formic acid and buffer B was 0.1% formic acid in 80% acetonitrile. A gradient of increasing organic proportion was used in combination with a reversed phase C18 separating column (2 µm particle size, 100 Å pore size, 15 cm length, 50 µm i.d., Thermo Fisher Scientific). Each MS scan was followed by a maximum of 10 MS/MS scans in the data-dependent mode. Blank samples were run between each set of three samples to minimize carry over.

The raw data files were analyzed with MaxQuant software (version 1.5.3.30) using Uniprot human FASTA database [49, 50]. Label-free quantification values (LFQ) and match between run options were selected. Intensity based absolute quantification (iBAQ) algorithm was also activated for subsequent relative protein abundance estimation [51]. The obtained protein files were analyzed by Perseus software (MQ package, version 1.6.12), in which contaminants and reverse hits were filtered out [50]. Protein identification based on non-unique peptides as well as proteins identified by only one peptide in the different
triplicates were excluded to increase protein prediction accuracy.

For identification of the bait interactors LFQ intensity-based values were transformed on the logarithmic scale (log2) to generate Gaussian distribution of the data. This allows for imputation of missing values based on the normal distribution of the overall data (in Perseus, width = 0.3; shift = 1.8). The normalized LFQ intensities were compared between grouped GFP triplicates and non-GFP triplicates using 1% as the permutation-based false discovery rate (FDR) in a two-tailed t-test. The threshold for significance (S0), based on the FDR and the ratio between GFP and non-GFP samples was kept at the constant value of 2. Relative plots were obtained by comparison of the iBAQ values of GFP interactors. The values of the non-GFP iBAQ values were subtracted from the corresponding proteins in the GFP pull-down and were next normalized on the menin-GFP bait protein for scaling and data representation purposes. All mass spectrometry data have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository under the dataset identifier PXD031928.

Genome localization experiments by (green)CUT&RUN

Genome localization analysis of GFP-tagged menin wild type, R52G, E255K, E359K or E408Q and GFP-tagged JunD was performed by greenCUT&RUN with the combination of enhancer-MNase and LaG16-MNase as described [33]. To localize MLL1 or RNA polymerase II standard CUT&RUN [32], was employed using MLL1/KMT2A antibodies (Epicypher #SKU 13-2004) or 8WG16 antibodies directed against the CTD of RPB1 at 1 µg/ml. For standard CUT&RUN and greenCUT&RUN mononucleosomal Drosophila DNA was used as spike-in DNA for normalization purposes and sequencing libraries were prepared as described [Nizamuddin et al., 2021]. In brief, purified DNA fragments were subjected to library preparation with NEB Next Ultra II and NEB Multiplex Oligo Set I/II as per manufacturer (New England Biolabs) protocol without size selection. For each library, DNA concentration was determined using a Qubit instrument (Invitrogen, USA) and size distribution was analyzed with Agilent Bioanalyzer chips (DNA high sensitivity assay). The 75-nucleotide paired-end sequencing reads were generated (Illumina, HiSeq 3000) with 6–32 M reads per sample (Additional file 6: Table S2). These NGS data have been deposited to Sequence Read Archive (SRA) portal of the NCBI under accession number PRJNA772915.

Bioinformatic analyses of genomics data

The HeLa cell datasets for H3K4me3 (ID: ENCCF063XTI), H3K4me1 (ENCCF617YCCQ) and H3K27ac (ENCCF113QJM) were downloaded from ENCODE. The ATACseq dataset for HeLa cells was downloaded from SRA-NCBI (Accession ID: SRR8171284). The datasets generated in house were initially passed through quality control using Trim-galore (version 0.6.3). Further reads were aligned on the human (version hg38) and Drosophila reference genome (BDFP5) using bowtie2 (version 2.3.4.1) with option: -dovetail -local -very-sensitive-local -no-unal -no-mixed -no-discordant -I 10 -X 700 [34, 53]. Equal number of reads was randomly selected for WT menin and menin mutants using sambamba (version 0.6.9) and utilized in the further analysis. The correlation plot was generated using the deetools (version 3.3.2) [54, 55].

Peaks were called using HOMER with default parameters except filtering based on clonal signals was disabled using option: -C 0. Both narrow and broad peaks were called with HOMER and merged together using bedtools (version 2.27.1) except for JunD for which only narrow peaks were called (Quinlan & Hall, 2010). Reads of the control samples were normalized with Spikeln and used to generate TagDirectories by HOMER using option: -totalReads before peak calling. To calculate normalized total reads, ratio of the Spikeln per human reads of control and experiment were calculated and multiplied with total number of control reads. In case of H3K4me3 (ENCFF862LUQ) the coordinate of the peaks were downloaded from ENCODE. The peaks of menin (wild type) overlapping with JunD, MLL1 and H3K4me3 were identified using bedtools and classified into eight categories using in-house script which can be downloaded [56].

HOMER was used to find AP1/ATF motifs within the peaks with default parameters. In differential peak analysis, menin peaks with coverage of fold changes ≤ 4 or p value ≥ 0.001 against mutants were considered as unaffected by the menin mutations.

Heatmaps were generated using deetools (version 3.3.2) with default parameters. All next-generation sequencing datasets have been deposited to the Sequence Read Archive (SRA) portal of the NCBI with accession ID PRJNA772915. The command-lines used for the data analysis can be downloaded [56].

Supplementary Information

The online version contains supplementary material available at https://doi.org/10.1186/s13072-022-00461-8.

Additional file 1: Figure S1. Quantification of menin mRNA and protein levels after transient transfections. Panel A. MEN1 mRNA expression was quantified by RT-qPCR and corrected for ACTB levels in pCDNA3.1 MEN1 WT and mutant transfected into HEK293 cells. The endogenous MEN1/ACTB mRNA ratios in the control pCDNA transfected cells were set at 1. The experiments were performed in triplicate. Error bars indicate SE. Panel B. Ratios of menin expression quantitation and MEN1 mRNA levels
The authors declare that they have no competing interest.

**Additional file 2: Figure S2.** Analysis of menin (mutant) proteins stably expressed in HeLa cells. Panel A. Fluorescence microscopy images of GFP-menin WT and mutant proteins stably expressed in HeLa cells stained with anti-GFP antibody. Zoomed regions were marked with white rectangles and are also provided in Fig. 2C. Panel B: Immunoblots of stable HeLa cell lines upon doxycycline for preparation of cytoplasmic and nuclear protein extracts. Expression of GFP-tagged menin was detected using GFP antibodies. α-tubulin and TBP antibodies were used as cytoplasmic and nuclear marker proteins and as loading controls. The position of co-migrated size markers are indicated on the left.

**Additional file 3: Table S2.** Summary of the data sets used in this study.

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**Author contributions**

KMAD, EOG, SN and HTMT wrote the manuscript. EJVDL, PVDH, JJK, KWR and MLB carried out the bioinformatics analysis of NGS data. All authors read and approved the final manuscript.

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**References**

1. Brandi ML, Agarwal SK, Perrier ND, Lines KE, Valk GD, Thakker RV. Multiple endocrine neoplasia type 1. Latest insights. Endocr Rev. 2020;42(2):1333.

2. Chandrasekharappa SC, Guru SC, Manickam P, Olufemi SE, Collins FS, Emmert-Buck MR, et al. Positional cloning of the gene for multiple endocrine neoplasia-type 1. Science. 1997;276(5311):404–7.

3. Jiao Y, Shi C, Edil BH, de Wilde RF, Klimstra DS, Maitra A, et al. DAXX/ATRX, MEN1, and mTOR pathway genes are frequently altered in pancreatic neuroendocrine tumors. Science. 2011;331(6021):199–203.

4. Hoadley KA, Yu C, Hinoue T, Wolf DM, Lazar AJ, Drill E, et al. Cell-of-origin patterns dominate the molecular classification of 10,000 tumors from 33 types of cancer. Cell. 2018;173(2):301–304.e6.

5. Dreijerink KMA, Timmers HTM, Brown M. Twenty years of menin: emerging opportunities for restoration of transcriptional regulation in MEN1. Endocr Relat Cancer. 2017;24(10):T135–45.

6. Hughes CM, Rozenblatt-Rosen O, Milea T, Copeland TD, Levine SS, Lee JC, et al. Menin associates with a trihexor family histone methyltransferase complex and with the hoxc8 locus. Mol Cell. 2004;13(4):587–97.

7. Lin W, Cao J, Liu J, Beshiri ML, Fujwara Y, Francis J, et al. Loss of the retinoblastoma binding protein 2 (RB2P2) histone demethylase suppresses tumorigenesis in mice lacking Rb1 or Men1. Proc Natl Acad Sci USA. 2011;108(33):13379–86.

8. Huang J, Gurung B, Wan B, Matkar S, Veniaminova NA, Wan K, et al. The same pocket in menin binds both MLL and JUND but has opposite effects on transcription. Nature. 2012;482(7366):542–6.

9. Agarwal SK, Guru SC, Heppner C, Erdos MR, Collins RM, Park SY, et al. Menin interacts with the API transcription factor JunD and represses JunD-activated transcription. Cell. 1999;96(1):143–52.

10. van Nuland R, Smits AH, Palka P, Jansen PW, Vermeulen M, Timmers HT. Quantitative dissection and stoichiometry determination of the human SET1/MLL histone methyltransferase complexes. Mol Cell Biol. 2013;33(10):2067–77.

11. Grembecka J, He S, Shi A, Purohit T, Muntean AG, Sorenson RJ, et al. Menin-MLL inhibitors reverse oncogenic activity of MLL fusion proteins in leukemia. Nat Chem Biol. 2012;8(3):277–84.

12. Ozery-Goeknar E, Nizamuddin S, Timmers HTM. A box of chemistry to inhibit the MEN1 tumor suppressor gene promoting leukemia. ChemMedChem. 2021;16(9):1391–402.

13. Yokoyama A, Somervaille TC, Smith KS, Rozenblatt-Rosen O, Meyerson M, Cleary ML. The menin tumor suppressor protein is an essential oncogenic cofactor for MLL-associated leukemogenesis. Cell. 2005;123(2):207–18.

14. Dzama MM, Steiner M, Rausch J, Sasca D, Schönfeld J, Kunz K, et al. Synergistic targeting of FLT3 mutations in AML via combined menin-MLL and FLT3 inhibition. Blood. 2020;136(21):2442–56.

15. Klossowski S, Miao H, Kemipinska K, Wu T, Purohit T, Kim E, et al. Menin inhibitor ML-3454 induces remission in MLL1-rearranged and NPM1-mutated models of leukemia. J Clin Invest. 2020;130(2):981–97.

16. Kvitnys AV, Evans K, Gadrey JY, Eschle BK, Hatton C, Uckelmann HJ, et al. A menin-MLL inhibitor induces specific chromatin changes and eradicates disease in models of MLL-rearranged leukemia. Cancer Cell. 2019;36(6):660–73.e11.

17. Uckelmann HJ, Kim SM, Wong EM, Hatton C, Giovannazzo H, Gadrey JY, et al. Therapeutic targeting of preleukemia cells in a mouse model of. Science. 2020;367(6477):386–90.

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