Thyrotropin receptor p.N432D retained variant is degraded through an alternative lysosomal/autophagosomal pathway and can be functionally rescued by chemical chaperones

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<td>Abstract:</td>
<td>Background. Loss of function mutations of thyrotropin receptor (TSHR) are one of the main causes of congenital hypothyroidism (CH). As for many disease-associated GPCRs, these mutations often affect the correct trafficking and maturation of the receptor, thus impairing the expression on the cell surface. Indeed, several retained GPCR mutants are able to effectively bind their ligands and to transduce signals when they are forced to the cell surface by degradation inhibition or by treatment with chaperones. Despite the large number of well-characterized retained TSHR mutants, no attempts have been made for rescue. Furthermore, little is known about TSHR degradation pathways. We hypothesize that, similarly to other GPCRs, TSHR retained mutants may be at least partially functional if their maturation and membrane expression is facilitated by chaperones or degradation inhibitors. Methods. We performed in silico predictions of the functionality of known TSHR variants and compared the results with available in vitro data. Western blot, confocal microscopy, ELISAs and dual luciferase assays</td>
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were used to investigate the effects of degradation pathways inhibition and of chemical chaperones treatments on TSHR variants maturation and functionality.

Results. Here, we report a high discordance rate between in silico predictions and in vitro data for retained TSHR variants, a fact indicative of a conserved potential to initiate signal transduction if these mutants were expressed on the cell surface. Indeed, we show for the first time experimentally that some maturation defective TSHR mutants are able to effectively transduce Gs/cAMP signaling if their maturation and expression are enhanced using chemical chaperones. Furthermore, through the characterization of the intracellular retained p.N432D variant, we provide new insights on TSHR degradation mechanism, as our results suggest that aggregation-prone mutant can be directed toward autophagosomal pathway instead of the canonical proteasome system.

Conclusions. Our study reveals alternative pathways for TSHR degradation. Retained TSHR variants can be functional when expressed on the cell surface membrane, thus opening the possibility of further studies on the pharmacological modulation of TSHR expression and functionality in patients in whom TSHR signaling is disrupted.
Thyrotropin receptor p.N432D retained variant is degraded through an alternative lysosomal/autophagosomal pathway and can be functionally rescued by chemical chaperones

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Abstract

**Background.** Loss of function mutations of thyrotropin receptor (TSHR) are one of the main causes of congenital hypothyroidism (CH). As for many disease-associated GPCRs, these mutations often affect the correct trafficking and maturation of the receptor, thus impairing the expression on the cell surface. Indeed, several retained GPCR mutants are able to effectively bind their ligands and to transduce signals when they are forced to the cell surface by degradation inhibition or by treatment with chaperones. Despite the large number of well-characterized retained TSHR mutants, no attempts have been made for rescue. Furthermore, little is known about TSHR degradation pathways. We hypothesize that, similarly to other GPCRs, TSHR retained mutants may be at least partially functional if their maturation and membrane expression is facilitated by chaperones or degradation inhibitors.

**Methods.** We performed *in silico* predictions of the functionality of known TSHR variants and compared the results with available *in vitro* data. Western blot, confocal microscopy, ELISAs and dual luciferase assays were used to investigate the effects of degradation pathways inhibition and of chemical chaperones treatments on TSHR variants maturation and functionality.

**Results.** Here, we report a high discordance rate between *in silico* predictions and *in vitro* data for retained TSHR variants, a fact indicative of a conserved potential to initiate signal transduction if these mutants were expressed on the cell surface. Indeed, we show for the first time experimentally that some maturation defective TSHR mutants are able to effectively transduce Gs/cAMP signaling if their maturation and expression are enhanced using chemical chaperones. Furthermore, through the characterization of the intracellular retained p.N432D variant, we provide new insights on TSHR degradation mechanism, as our results suggest that aggregation-prone mutant can be directed toward autophagosomal pathway instead of the canonical proteasome system.
Conclusions. Our study reveals alternative pathways for TSHR degradation. Retained TSHR variants can be functional when expressed on the cell surface membrane, thus opening the possibility of further studies on the pharmacological modulation of TSHR expression and functionality in patients in whom TSHR signaling is disrupted.
Introduction

Loss of function (LOF) mutations of thyrotropin receptor (TSHR) are one of the principal causes of congenital hypothyroidism (1, 2). TSHR is a G-protein coupled receptor (GPCR) characterized by a seven transmembrane alpha-helix B-subunit and an extracellular A-subunit, linked by disulphide bonds (3, 4).

TSHR post-translational modifications are required for correct trafficking, maturation, and activity (3, 5, 6). The N-linked glycosylation happening in the endoplasmic reticulum (ER) is fundamental (Fig. 1A): acquisition of mannose-type carbohydrates permits the interactions with molecular chaperones (Fig. 1B), required for correct receptor folding, homodimerization, passage through ER quality control system and translocation to cis-Golgi (Figure 1C) (3, 5, 7). In trans-Golgi, TSHR finally acquires the complex-type carbohydrates (Fig. 1D) that characterize the mature form expressed on cell surface and undergoes tyrosine sulfation, fundamental for high-affinity binding and activation (6, 8). On plasma membrane TSHR is cleaved by an unidentified enzyme with loss of a short sequence of variable size, peptide C (Figure 1E). The receptor is finally composed by an extracellular A-subunit and a transmembrane B-subunit linked by disulfide bonds (3, 5, 9, 10).

For many disease-associated GPCRs, including TSHR, LOF is most often due to poor cell surface expression, rather than from intrinsic deficiencies in signal transduction. The abnormal mutant conformation leads to interactions with alternative molecular chaperones (3, 11, 12), ER blockage and degradation by proteasome or by autophagosome (Fig. 1G, H) (13, 14). Different retained GPCR mutants are able to effectively bind their ligands and transduce intracellular signals when forced to cell surface (15–17). The use of chemical chaperones is a well explored area to overcome ER retention of various membrane receptors (18–20). Nowadays, little is known about TSHR degradation pathways and no attempts have been made in TSHR mutants’ rescue.
The aim of our work is to better elucidate TSHR degradation pathways and the possibility of TSHR mutants rescue with chemical chaperones. We concentrated our attention on two different mutants that we previously described: the TSHR p.N432D, which is retained in the ER as high-mannose form, and the p.P449L, that is normally expressed on plasma membrane but with impaired signaling (21). We then validate our findings in two other retained TSHR mutants (22, 23).

Our results show for the first time that maturation-defective TSHR mutants are able to transduce Gs/cAMP signaling when rescued by the chemical chaperone Trimethylamine-N-oxide (TMAO). Moreover, we provide new insights on TSHR degradation mechanism, as our results suggest that aggregation-prone mutants are directed toward the autophagosomal pathway instead of the canonical proteasome system.

Materials and Methods

Chemicals

Cell culture reagents, ProLong Gold Antifade Reagent with DAPI, LysoTracker Red DND-99, ER-Tracker Green, Alexa-Fluor conjugated and HRP-conjugated antibodies, Restore Western Blot Stripping reagent were purchased from Thermo-Fisher. Mouse Anti-Actin Ab-5 was purchased from BD Biosciences. Anti TSHR antibodies BA8 (Cat#SC_BA8, RRID:AB_2716681), 3G4 (Cat#SC_3G4, RRID:AB_2716682) and 28.1 (Cat#SC_28.1, RRID:AB_2716683) were described elsewhere (24–28) and were a kind gift from Dr S. Costagliola (IRIBHM, ULB, Brussels). Anti E-Cadherin antibody was purchased from Abcam, anti VDAC was purchased from Santa Cruz. bTSH, Anti-GFP antibody, TMAO, DMSO and MTT were purchased from Sigma-Aldrich.
**In silico prediction**

TSHR variants membrane expression and functionality was assessed through the TSH receptor mutation database (29). 55 variants were subjected to *in silico* predictions and assigned as damaging or not damaging as specified in Supplementary Methods.

**Cell culture, transfection, treatments and viability assay**

COS-7 cells were grown in DMEM (Gibco) supplemented with 10% fetal bovine serum (Sigma-Aldrich) and penicillin-streptomycin (Sigma-Aldrich). TSHR cloning and mutagenesis were described elsewhere (21). pSVL plasmids containing WT, p.E34K and p.R46P TSHR variants were a kind gift of Dr. Tonacchera (22, 23).

Transfection, degradation modulation and rescue were performed as described in Supplementary Methods. Cell viability was tested with MTT assay (30).

**Western blotting**

Cells were lysed in SDS buffer (62.5 mM Tris-HCl pH 6.8, 2% sodium dodecyl sulfate) supplemented with protease, phosphatase and proteasome inhibitors.

Membrane preparations were obtained with Plasma Membrane Protein Extraction Kit (Abcam) following manufacturer’s instructions. Total Cellular Membranes and Plasma Membranes fractions were then processed as the other samples, as described in Supplementary Methods.

Band intensity was quantified with ImageJ software (31).

**Immunofluorescence and Confocal Microscopy**
Samples were processed as previously described (32), and detailed in Supplementary Methods.

Images were acquired with EclipseTi-E inverted microscope with IMA10X Argon-ion laser System by Melles Griot; images were acquired with CFI Plan Apo VC 60X Oil (Nikon).

Flow cytometry

Samples were processed as previously described (32), and detailed in Supplementary Methods. Measurements were performed with FACSCalibur flow cytometer (Becton Dickinson). Data were analyzed with Flowing Software 2.

Functional assays

cAMP pathway activity was assessed with Cignal CRE Reporter (luc) Kit (Quiagen), while Gq11/IP3 pathway activity was measured with IP-One ELISA assay kit (Cisbio), following manufacturer’s instructions, as described in Supplementary Methods.

Statistical analysis

All experiments were independently repeated at least three times, as indicated in the figure legends. After normal distribution and variance similarity evaluation, two-sided unpaired t-test (eventual Welch’s correction for groups with different variances), one-way ANOVA with Bonferroni post-hoc test, Kruskal-Wallis H test with Dunns post hoc test and Chi-square test were used as indicated in figures’ legend.
For concentration-effect curves of Gs/cAMP signaling a log(agonist) vs. normalized response -
Variable slope equation was used for curve interpolation and parameters definition.

For confocal experiments, the degree of colocalization was quantified through Pearson’s correlation coefficient, as measured with Nikon NIS Elements software. Correlation was defined as strong with Pearson’s correlation coefficient bigger than 0.8, moderate when bigger than 0.5 and weak when bigger than 0.2.

In all figures data are shown as mean±SEM, analyzed using GraphPad Prism 5 software and significance expressed as P values (* p < 0.05, ** p< 0.01, *** p < 0.001).

Results

In silico prediction and in vitro data of receptor functionality are significantly discordant in retained mutants

We obtained complete information about in vitro functionality and subcellular localization of 55 LOF TSHR variants (29) and categorize them as intracellular-retained or membrane-expressed (Suppl. Table 1).

These mutations were subjected to in silico predictions and assigned as functional or non-functional. The comparison of in vitro and in silico data reveals significantly higher discordance rate among the retained group (12/24 and 7/31 mutants with positive prediction but in vitro LOF for intracellular-retained and membrane-expressed respectively, p=0.0471) (Table 1).

Such discrepancy may indicate that some ER retained mutants can potentially transduce signal if expressed on cell surface.
We thus explored the degradation mechanisms and chaperone rescue on two previously reported (21) TSHR LOF variants: the intracellular-retained p.N432D and the membrane-expressed p.P449L.

N432D variant is arrested in the ER and forms different aggregates

We performed confocal microscopy with two different antibodies, the BA8 directed against a conformational epitope on the mature A-subunit and the 3G4 raised against a linear epitope in the C-peptide that recognizes principally immature forms (24, 25).

WT TSHR and p.P449L variant have a normal membrane expression in all transfected cells (Fig. 2 A), whereas p.N432D have a variable pattern detected by BA8 antibody, with three main morphologies: small intracellular aggregates (SA), perinuclear signal (PS) and cytoplasmic macroaggregate (MA) (Fig. 2 B). SA and PS are the most frequent ones, while in around 10% p.N432D pattern has the characteristics of more than one morphology (mixed morphology, MM) (Fig. 2 C). In contrast, p.N432D variant staining with 3G4 antibody reveals a constant pattern of diffuse perinuclear signal that is not detectable with BA8 antibody (Fig. 2 B, second lane). This difference may indicate the presence of a significant amount of immature or incorrectly folded receptors recognized only by 3G4 antibody but not by BA8 (24).

Interestingly, SA are similar to the puncta that characterize misfolded GPCR mutants degraded by autophagocytosis (33, 34), while MA are suggestive of perinuclear aggregates related to the proteasome degradation pathway (34, 35).

Co-staining with p.N432D variant and ER or late endosome/lysosome markers shows that the majority of the protein recognized by 3G4 antibody is indeed localized in the ER (Fig. 2 D, E).

Different features of ER stress, like vacuoles and enlarged morphology (36), are also detected in
transfected cells. On the other hand, the aggregates recognized by BA8 antibody show a mild co-localization with endosomes/lysosomes (Fig. 2 F, G).

TSHR mutants are degraded through different pathways

For many GPCRs, ubiquitin-proteasome system is the main degradation system (37, 38), while mutants prone to form aggregates are directed toward autophagic degradation (33, 39). We evaluated if that was our case by performing western blot in different conditions, with the 28.1 antibody that recognizes full length receptor at different stages of maturation together with the cleaved A-subunit (28, 40).

MG132 proteasome inhibitor induces a significant accumulation of mature WT TSHR and p.P449L variant, confirming the fundamental role of this pathway. However, only a strong accumulation of the immature form is detected for p.N432D (Fig. 3 A, B). NH₄Cl autophagy inhibition does not cause significant alterations in the total WT TSHR, although a change in the amount of mature forms can be appreciated, as previously reported (5). On the other hand, endolysosomal inhibition induces a more effective accumulation of p.P449L and of immature p.N432D than the proteasomal one (Fig. 3 A, B). Confocal microscopy experiments reveal significant increase in SA after autophagy inhibition, while a significant increase in MA is seen after proteasome inhibition; concomitant inhibition has indeed an intermediate effect (Fig. 3 C), thus confirming western blot data.

Autophagy activation with LiCl induces an almost complete degradation of p.N432D variant, with milder effects on p.P449L and no effects on WT (Fig. 3 D, E). Moreover, only p.N432D expression induces JNK 1/2 phosphorylation, an event linked to autophagy activation (41), and significantly reduces cell viability (Suppl. Fig. 1 A, B), thus confirming the role of autophagy in misfolded TSHR degradation.
The chemical chaperone TMAO restores p.N432D mutant membrane expression.

p.N432D mutant does not maturate even if protein degradation is inhibited, but as immature TSHR can signal when expressed on plasma membrane (42, 43), we investigated whether this was our case by the use of different chemical chaperones.

Western blot experiments show that, unlike in other GPCRs (44), treatment with glycerol is not effective in p.N432D variant rescuing (Suppl. Fig. 3). Nevertheless, TMAO treatment (45) increases the maturation of all TSHR variants, but with major effects on p.N432D whose A subunit intensity reaches levels similar to WT control indicating possible membrane expression, while the high mannose form has a larger increase in the WT and p.P449L (Fig. 4 A, B). The hybridization with 3G4 antibody reveals that TMAO treatment causes a significant increase in a high molecular weight band (around 200 kDa) (Suppl. Fig. 2 A, B) which has been identified as dimers of high mannose forms (43), whose formation is fundamental for passage through ER quality control.

FACS experiments in non-permeabilized cells confirmed cell-surface expression of TMAO treated p.N432D. Interestingly, membrane expression resulted 75% of WT control with BA8 antibody (Fig. 4 C, D), and only around 50% of WT control with 3G4 antibody (Fig. 4 E, F) with a BA8:3G4 ratio of 1.62±0.25 (p<0.05 vs treated WT), a finding that indicates increased cleavage of the p.N432D mutant (25).

Immunofluorescence experiments with BA8 staining confirm p.N432D membrane expression after TMAO treatment. The increase in intracellular staining for TMAO-treated TSHRs is in agreement with the increase in immature forms detected at western blot experiments (Fig. 4 G).

Cellular membranes fractionations revealed that the A subunit is the predominant form on the cell surface. Moreover, TMAO promotes WT TSHR translocation on the cell surface, as we detected a
decrease in the levels of all TSHR maturation forms in the total membrane extracts and an increase in the plasma membrane extracts. This effect is not seen in the p.P449L variant, that is also insensible to TMAO effects (Suppl. Fig. 2 C, D).

**Membrane expression uncovers the functional potential of p.N432D and other retained variants.**

The evaluation of rescued p.N432D variant signaling transduction abilities through Gs/cAMP and Gq11/IP3 pathways revealed that indeed the mutant is partially functional when expressed on plasma membrane. In fact, although Gq11/IP3 pathway remains greatly compromised (Fig. 5 A), the maximal Gs/cAMP response is almost completely rescued (Fig. 5 B). Concentration-effect curves (Fig. 5 C, Suppl. Fig. 4 A and Table 2) show that TMAO treatment has virtually no effect on the Gs/cAMP signaling of either the WT or p.P449L mutant receptor, while the TMAO-treated p.N432D curve is right-shifted, indicating higher EC50 values.

As last step, we investigated two additional retained variants that have discordant *in silico* and *in vitro* functionality: the p.E34K, that has a reported membrane expression of 30% of WT, and the p.R46P that is reported to be almost totally retained and with very low ability to signal through the cAMP pathway (22, 23).

TMAO treatment induces an increase in the cleaved A-subunit levels in the p.E34K variant and greatly enhances the maturation of the retained p.R46P one (Fig. 5 E, D), with effects similar to the ones observed in p.N432D. Accordingly, functional assays reveal a significant increase in both Gs/cAMP and Gq11/IP3 pathways for p.E34K and a significant rescue of the signaling abilities of p.R46P (Fig. 5 F, G), with concentration-effect curves and EC50 similar to those of the WT (Fig. 5 H).
Discussion

In the present work, we reveal two important issues regarding the possible intracellular destiny of the folding-defective TSHR mutants. First, they may be degraded not only through the proteasomal pathway, but also through an alternative autophagosomal-like pathway that kicks in as emergency exit after retention in the ER. Second, they can at least partially function if forced to the cell surface by using chemical chaperones. Our data provide a possible explanation for the observed lack of concordance between in silico prediction of receptor functionality and in vitro findings, as misfolded mutants that retain signaling abilities may have a premature maturation arrest, intracellular retention and subsequent degradation.

The involvement of the lysosomal system in the degradation of misfolded TSHR mutants is a new interesting finding. In particular, p.N432D has such structural changes that prevent passing the ER quality control. In the ER the mutant is likely to form aggregates, microscopically detected as SA pattern, that cannot be retro-translocated to the cytoplasm where proteasome operates, but are instead degraded by alternative autophagocytosis (Fig.1, 2, 3 and Suppl. Fig. 1); a behavior similar to the one previously described for gonadotropin releasing-hormone receptor (GnRHR) mutant p.E90K (33).

The treatment with TMAO is likely inhibiting the formation of ER aggregates while promoting the receptor homodimerization, sheltering p.N432D from ER quality control and allowing advancement to Golgi compartment and finally to plasma membrane (3, 46), as also indicated by the appearance of the A-subunit bands in plasma membrane preparations. Nevertheless, its maturation does not seem to follow the regular steps even after TMAO treatment as we detected very low levels of complex carbohydrates form (Fig. 4A and 4B).

There are two possible explanations of this issue. The first and most likely hypothesis is that only a small percentage of plasma membrane p.N432D mutant reach full maturation, while most of it
still blocked at the high-mannose stage. Membrane expression of immature TSHR has already been described (26, 42, 43, 47) and TSHR with reduced glycosylation sites has TSH binding affinity and EC50 for cAMP that are indistinguishable from the mature one (3). In this case, the p.N432D maturation limiting factor may be the ability to form dimers in the ER compartment, as the staining with 3G4 antibody promptly showed a significant increase in the levels of immature TSHR dimers after TMAO treatment (Suppl. Fig. 2 A, B). The increased cleavage indicated by the variation in BA8:3G4 ratio (Fig. 4 C, E, Table 2) can then be explained by the already known higher sensitivity to proteases action of immature TSHR (25, 42).

The second possible explanation is that TMAO-treated p.N432D mutant reaches full maturation, but all the mature receptor undergoes proteolytic cleavage and thus only A-subunit is detected. This may be explained by an increased sensitivity of the TSHR mutant to proteases or because a lower amount of mutant TSHR on the membrane more effectively processed by proteases.

Irrespective of these considerations, functional assays show that p.N432D mutant is able to bind TSH and transduce intracellular signal when expressed on plasma membrane (Fig. 5 A-C and Table 2). The lack of Gq11/IP3 pathway activity may be explained by the intrinsic differences between Gs and Gq interaction with TSHR. First of all, the Gq11/IP3 pathway is more dependent upon the total amount of cleaved receptor (10) and on TSHR homodimerization abilities (48) and TMAO treated p.N432D has an absolute amount of cleaved receptor present on plasma membrane definitely lower than WT one (Fig. 4 and Suppl. Fig. 2). In addition, interactions between TSHR and Gq are more demanding than the ones with Gs (4, 49), and an in silico model predicted that p.N432D mutation severe modifications can affect the interaction with G-protein (21). The treatment with TMAO can either mask these conformational alterations or force the mutant through a more correct conformation that is enough to achieve a partial rescue of Gs interactions and cAMP signaling but is not enough to restore the more demanding interactions with Gq.
These speculations are also supported by the findings on two other discordant mutants, where TMAO treatment more efficiently rescues Gs/cAMP than the Gq11/IP3 signaling (Fig. 5 D-H and Suppl. Fig. 4 B).

In conclusion, our work shows that TSHR can be degraded through proteasome or autophagosome pathways depending on specific structural defects. The chaperone TMAO allows TSHR mutants to pass ER quality control, increasing cell surface expression. As for other GPCR-related diseases, TSHR LOF mutations are mainly causing ER retention, as detected by the discrepancy between in silico predictions and in vitro data. As we demonstrated here, retained mutants that are brought to the cell surface are able to effectively transduce intracellular signal. These findings open the possibility of further studies on pharmacological modulation of TSHR expression and functionality in patients with disrupted TSHR signaling.
Acknowledgments

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Grassi ES, Labadi A and Ghiandai V designed and performed the experiments.

Vezzoli V and Bonomi M contributed to the experiment planning.

LP supervised the experimental work and provided research funds.

All authors contributed to the writing and revision of the manuscript.

Author Disclosure Statement

No competing financial interests exist.
References


Grassi ES, Vezzoli V, Negri I, et al. SP600125 has a remarkable anticancer potential against
undifferentiated thyroid cancer through selective action on ROCK and p53 pathways.


44. Chen DN, Ma YT, Liu H et al. Functional rescue of Kallmann syndrome-associated


Table 1: *in vitro* and *in silico* functionality concordance of TSHR variant with different subcellular localization. Distribution of TSHR variants from *in vitro* data and *in silico* predictions concordance of function in relation to subcellular localization. For 55 different variants data about *in vitro* subcellular localization and functionality were obtained through literature review. *In silico* prediction were obtained with 6 different online tools and each variant was then assigned to one of the four groups: membrane localization with *in vitro* and *in silico* concordance on functionality, membrane localization with *in vitro* and *in silico* discordance on functionality, intracellular retainment with *in vitro* and *in silico* concordance on functionality, intracellular retainment with *in vitro* and *in silico* discordance on functionality.

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Table 2: membrane expression and functional parameters of the LOF TSHR variants. The table summarize the main characteristics of the LOF TSHR variants. Membrane expression was examined with flow cytometry as in figure 4C and 4E. Maximal stimulation and EC50 were obtained from experiment as in Figure 5. Values are expressed as mean±SD.

Statistical analysis: statistical significance was determined with One-Way ANOVA. * p<0.05, ** p<0.01, ***p<0.001 vs. WT TSHR, §§§ p<0.001 vs. the respective untreated TSHR variant.

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<th>Membrane expression (3G4)</th>
<th>Membrane expression (BA8)</th>
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<th>cAMP Emax (% WT)</th>
<th>cAMP EC50 (U/L)</th>
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<td>WT</td>
<td>100.00 ± 1.09</td>
<td>100.00 ± 0.61</td>
<td>100.00 ± 10.41</td>
<td>100.00 ± 1.80</td>
<td>0.10 ± 0.04</td>
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<td>WT + TMAO</td>
<td>102.90 ± 1.31</td>
<td>97.17 ± 2.75</td>
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<td>101.00 ± 6.33</td>
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<td>p.N432D</td>
<td>1.75 ± 2.06</td>
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<td>9.12 ± 2.08</td>
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<td>p.N342D + TMAO</td>
<td>51.03 ± 9.19</td>
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For Peer Review ONLY/ Not for Distribution
Figure legends.

**Figure 1: TSHR maturation and the hypothesized degradation pathways.**

TSHR is constituted of a seven-transmembrane alpha helix B-subunit (green) and an extracellular A-subunit (purple). It is synthesized in the endoplasmic reticulum as a full length unglycosylated protein (A, immature form). The acquisition of mannose-type carbohydrates (B, high-mannose form) permits the interactions with molecular chaperones like calnexin and calreticulin, thus facilitating protein folding and is required for receptor homodimerization, allowing passage through ER quality control system and translocation to cis-Golgi (F). The TSHR maturation continues (C) as it passes to the trans-Golgi where it acquires the complex-type carbohydrates that characterize the mature form of the protein (D, complex carbohydrates form) that is expressed on the cell surface. The majority of mature TSHR present on cell membrane is cleaved with loss of a small aminoacidic sequence of variable size, named peptide C. The receptor is thus finally composed of a transmembrane B-subunit linked by disulfide bonds to the extracellular A-subunit that can have slightly different dimensions depending on the cleavage sites and carries all the carbohydrate side chains (E, grey discontinued line indicates that only A-subunit is shown in western blot image).

The TSHR can be degraded through different systems. The misfolded receptors not able to proceed through the endoplasmic reticulum may be directed toward the proteasome (G) or the endolysosome (H) systems, depending on the nature of the alterations. Receptors not able to proceed through maturation in the Golgi can be retrotranslocated to the ER (F) for another round of control or directed toward degradation (I). The turnover of normal membrane TSHR also occurs through the endolysosome system after receptor endocytosis (I).
Figure 2: p.N432D mutant is retained in the ER and lysosomes in different aggregates.

A: representative images of WT TSHR, p.P449L mutant and empty vector transfected cells stained with 3G4 and BA8 antibodies.

B: representative images of the different morphological presentations of p.N432D mutant after anti-TSHR 3G4 or anti-TSHR BA8 antibodies staining. For BA8 antibody: PS, perinuclear signal; SA, small intracellular aggregates; MA, macroaggregates; MM, mixed morphology, concomitant presence of PS, SA and/or MA detected with BA8 antibody. 3G4 antibody detected a constant mix of perinuclear signal and small aggregates.

C: quantification of the relative frequencies of the different p.N432D patterns detected with BA8 antibody as in Fig. 2B. PS, perinuclear signal; SA, small intracellular aggregates; MA, macroaggregates; MM, mixed morphology, concomitant presence of PS, SA and/or MA detected with BA8 antibody.

D: representative images of colocalization experiment showing endoplasmic reticulum (ER, red) and N432D mutant (green) stained with either BA8 or 3G4 antibodies.

E: analysis of Pearson’s coefficient for colocalization. The graph represents the averages of the Pearson’s coefficients for colocalization detected for N432D mutant stained with either BA8 or 3G4 antibodies and Endoplasmic Reticulum.

F: representative images of colocalization experiments showing late endosome/lysosomes (LYSO, red) and N432D mutant (green) stained with either BA8 or 3G4 antibodies.

G: analysis of Pearson’s coefficient for colocalization. The graph represents the averages of the Pearson’s coefficients for colocalization detected for N432D mutant stained with either BA8 or 3G4 antibodies and late endosome/lysosomes.
Statistical analysis: C n=14 (2018 cells analyzed), E n=12 (248 cells analyzed), G n=6 (110 cells analyzed). Statistical significance was determined with One-Way ANOVA in C and t-test with Welch’s correction in E and G. * p<0.05 and ***p<0.001 as indicated.

Figure 3: WT and mutant TSHRs are degraded through different pathways.

A: representative western blot images of TSHR expression and maturation after treatment with 20 mM NH₄Cl, 10 µM MG132 and a combination of the two inhibitors. TSHR was stained with 28.1 antibody, GFP was used as transfection efficiency control and actin was used as loading control.

B: densitometric quantification of western blot experiments showing complex carbohydrates, high mannose and A subunit forms of TSHR after treatment with NH₄Cl, MG132 and a combination of the two inhibitors.

C: representative images and relative quantification of confocal microscopy experiments showing anti-TSHR BA8 antibody staining of p.N432D mutant after treatment with NH₄Cl, MG132 and a combination of the two inhibitors. For each treatment, the signal pattern with bigger fold change increase in respect to control is shown. White, perinuclear signal (PS); light grey, small intracellular aggregates (SA); intermediate grey, macroaggregates (MA); dark grey, mixed morphology (MM).

D: representative western blot images of TSHR expression and maturation after treatment with 10 mM LiCl. TSHR was stained with 28.1 antibody, GFP was used as transfection efficiency control and actin was used as loading control.

E: densitometric quantification of western blot experiments showing complex carbohydrates, high mannose and A subunit forms of TSHR after treatment with LiCl.

Statistical analysis: B n=4, C n=14, E n=3. Statistical significance was determined with One-Way ANOVA (non-parametric Kruskal-Wallis H test) followed by Dunn’s post hoc test.
*p<0.05, ** p<0.01 and ***p<0.001 as indicated.
Figure 4: TMAO restores p.N432D mutant trafficking and membrane expression.

A: representative images of western blotting experiments showing the expression and maturation of the TSHR variants without and after treatment with TMAO. TSHR was stained with 28.1 antibody, GFP was used as transfection efficiency control and actin was used as loading control.

B: densitometric quantification of western blot experiments showing complex carbohydrates, high mannose and A subunit forms of TSHR without and after TMAO treatment.

C: representative flow cytometric histograms showing BA8 antibody signal in unpermeabilized cells with or without TMAO treatment. The R-1 markers indicate cells expressing TSHR and were used to quantitate receptor expression based on the mean fluorescence intensity. Untreated samples, light blue area; TMAO-treated samples, red area; overlapping area, green.

D: mean fluorescence intensity quantification of the TSHR variants expression on cell membrane without or after TMAO treatment, with 3G4 antibody labeling. Values are expressed as percentage of untreated WT.

E: representative flow cytometric histograms showing 3G4 antibody signal in unpermeabilized cells with or without TMAO treatment. The R-1 markers indicate cells expressing TSHR and were used to quantitate receptor expression based on the mean fluorescence intensity. Untreated samples, light blue area; TMAO-treated samples, red area; overlapping area, green.

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G: representative images of confocal microscopy experiments performed with BA8 staining, showing TSHR variants in normal conditions and after TMAO treatment. TSHR, green staining, nuclei, blue DAPI staining.
Statistical analysis: B n=3, D n=6, F n=6. Statistical significance was determined with One-Way ANOVA followed by Dunns (D,F) or Bonferroni (B) post hoc test.

*p<0.05, **p<0.01, ***p<0.001 in respect to untreated TSHR WT.
Figure 5: TMAO treatment unveiled partial functionality of N432D and other known retained mutants.

A: Gq11/IP3 pathway activity after maximal dose stimulation of TSHR WT, p.N432D and p.P449L variants upon normal conditions and after TMAO treatment, measured as IP1 accumulation and expressed as percentage of stimulated WT activity.

B: Gs/cAMP pathway activity after maximal dose stimulation of TSHR WT, p.N432D and p.P449L variants upon normal conditions and after TMAO treatment, measured as cAMP reporter luciferase activity and expressed as percentage of stimulated WT activity.


D: representative images of western blotting experiments showing the expression and maturation of the TSHR variants without and after treatment with TMAO. TSHR was stained with 28.1 antibody, actin was used as loading control.

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G: Gs/cAMP pathway activity after maximal dose stimulation of TSHR WT, p.E34K and p.R46P variants upon normal conditions and after TMAO treatment, measured as cAMP reporter luciferase activity and expressed as percentage of stimulated WT activity.

Statistical analysis: A, B, C n=8, E, F, G, H n=3. Statistical significance was determined with One-Way ANOVA followed by Bonferroni’s post hoc test (A, B, C) or Kruskal-Wallis H test followed by Dunn’s post hoc test (E, F, G).

*p<0.05, **p<0.01, ***p<0.001 in respect to TMAO untreated TSHR WT or as indicated.
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B: densitometric quantification of western blot experiments showing complex carbohydrates, high mannose and A subunit forms of TSHR after treatment with NH4Cl, MG132 and a combination of the two inhibitors.  
C: representative images and relative quantification of confocal microscopy experiments showing anti-TSHR BA8 antibody staining of p.N432D mutant after treatment with NH4Cl, MG132 and a combination of the two inhibitors. For each treatment, the signal pattern with bigger fold change increase in respect to control is shown. White, perinuclear signal (PS); light grey, small intracellular aggregates (SA); intermediate grey, macroaggregates (MA); dark grey, mixed morphology (MM).  
D: representative western blot images of TSHR expression and maturation after treatment with XX mM LiCl. TSHR was stained with 28.1 antibody, GFP was used as transfection efficiency control and actin was used as loading control.
E: densitometric quantification of western blot experiments showing complex carbohydrates, high mannose and A subunit forms of TSHR after treatment with LiCl. Statistical analysis: B n=4, C n=14, E n=3. Statistical significance was determined with One-Way ANOVA (non-parametric Kruskal-Wallis H test) followed by Dunn’s post hoc test. *p<0.05, ** p<0.01 and ***p<0.001 as indicated.
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Figure S5: TMAO treatment unveiled partial functionality of N432D and other known retained mutants.

A: Gq11/IP3 pathway activity after maximal dose stimulation of TSHR WT, p.N432D and p.P449L variants upon normal conditions and after TMAO treatment, measured as IP1 accumulation and expressed as percentage of stimulated WT activity.

B: Gs/cAMP pathway activity after maximal dose stimulation of TSHR WT, p.N432D and p.P449L variants upon normal conditions and after TMAO treatment, measured as cAMP reporter luciferase activity and expressed as percentage of stimulated WT activity.


D: representative images of western blotting experiments showing the expression and maturation of the TSHR variants without and after treatment with TMAO. TSHR was stained with 28.1 antibody, actin was used as loading control.

F: Gq11/IP3 pathway activity after maximal dose stimulation of TSHR WT, p.E34K and p.R46P variants upon normal conditions and after TMAO treatment, measured as IP1 accumulation and expressed as percentage of stimulated WT activity.

G: Gs/cAMP pathway activity after maximal dose stimulation of TSHR WT, p.E34K and p.R46P variants upon normal conditions and after TMAO treatment, measured as cAMP reporter luciferase activity and expressed as percentage of stimulated WT activity.


Statistical analysis: A, B, C n=8, E, F, G, H n=3. Statistical significance was determined with One-Way ANOVA followed by Bonferroni’s post hoc test (A, B, C) or Kruskal-Wallis H test followed by Dunn’s post hoc test (E, F, G).

*p<0.05, **p<0.01, ***p<0.001 in respect to TMAO untreated TSHR WT or as indicated.
Supplementary Methods

Chemicals

Cell culture reagents, ProLong Gold Antifade Reagent with DAPI, LysoTracker Red DND-99, ER-Tracker Green, Alexa-Fluor conjugated and HRP conjugated secondary antibodies, Restore Western Blot Stripping reagent were purchased from Thermo Fisher. Purified Mouse Anti-Actin Ab-5 was purchased from BD Biosciences. Anti TSHR antibodies BA8 (Cat# SC_BA8, RRID:AB_2716681), 3G4 (Cat# SC_3G4, RRID:AB_2716682) and 28.1 (Cat# SC_28.1, RRID:AB_2716683) were described elsewhere (1–5) and were a kind gift from Dr S. Costagliola (IRIBHM, ULB, Brussels). Anti E-Cadherin antibody was purchased from Abcam, anti VDAC was purchased from Santa Cruz. bTSH, Anti-GFP antibody, Trimethylamine-N-oxide (TMAO), DMSO and MTT were purchased from Sigma-Aldrich.

In silico prediction

The membrane expression and functionality of TSHR variants was assessed through the TSH receptor mutation database (6) at http://endokrinologie.uniklinikum-leipzig.de/tsh/frame.html and subsequent extensive literature research of references provided in the database. We obtained complete information for 55 variants that were then subjected to in silico predictions with 6 different tools: polyphen-2 (7) (genetics.bwh.harvard.edu/pph2/), PROVEAN (8) (provean.jcvi.org/), SIFT (9) (sift.jcvi.org/www/SIFT_enst_submit.html), PhD-SNP (10) (snps.biofold.org/phd-snp/phd-snp.html), PANTHER (11) (www.pantherdb.org/tools/csnp), SNPs&GO (12) (snps.biofold.org/snps-and-go/snps-and-go.html). The TSHR variants were then assigned as damaging (more than 50% of predictions concordant as functionally damaging or impaired functionality) or not damaging (50% or more of predictions concordant as neutral or benign).
Cell culture, transfection, treatments and viability assay

COS-7 cells were grown in DMEM medium (Gibco) supplemented with 10% fetal bovine serum (Sigma Aldrich) and penicillin-streptomycin mixture (Sigma Aldrich). TSHR cloning and mutagenesis were described elsewhere (13). In cells transfected with pIRES2-EGFP vector, EGFP is an optimal internal normalizer of the transfection efficiency, without interfering with the TSHR maturation and expression.

pSVL plasmids containing WT, p.E34K and p.R46P TSHR variants were a kind gift of Dr. Tonacchera and have been previously described (14, 15). For all the experiments, 3.5x10⁵ cells per well were seeded in 6-well plates in order to obtain similar confluency in all the conditions, as confluency directly influences TSHR cleavage (16). After 24 hours, in each well 1µg of plasmid DNA was transiently transfected with Fugene Transfection Reagent (Promega) following the manufacturer’s instructions. All samples were then analyzed 48 hours after transfection. For degradation inhibition or induction cells were treated with 10 µM MG132, 20 mM NH₄Cl or 10 mM LiCl 24 hours after transfection up until sample analysis. For chemical chaperone rescue, cells were cultured in medium containing 10% glycerol or 100 mM TMAO from six hours after transfection up until sample analysis. Cell viability was tested at the indicated times with MTT assay, as previously described (17).

Western blotting

Cells were lysed in SDS sample buffer (62.5 mM Tris-HCl pH 6.8, 2% sodium dodecyl sulfate) supplemented with protease, phosphatase and proteasome inhibitors.

Membrane preparations were obtained with Plasma Membrane Protein Extraction Kit (Abcam) following manufacturer’s instructions. Total Cellular Membranes and Plasma Membranes fractions
were then processed as the other samples. The samples were heated for 3 min at 95°C and sonicated. 40 µg of protein extracts were then separated on NuPage 4-12% Bis-Tris Gels (Thermo fisher) and transferred with iBlot System (Thermo fisher). Membranes were blocked with 5% nonfat dry milk in TBS-T solution for 1 hour at room temperature and probed overnight at 4°C with monoclonal anti-TSHR antibody (clone 28.1) (4, 5) used as hybridoma supernatant and diluted to 1:20 ratio in the blocking buffer. Actin, E-cadherine and VDAC were used as loading control, GFP as transfection efficiency control. After washing, membranes were incubated for 1 h at room temperature in the presence of horseradish peroxidase-conjugated goat anti-mouse IgG secondary antibody (Merck Millipore) diluted to 1:10000 ratio in blocking buffer. Detection was performed utilizing Luminata Forte ECL (Merck Millipore).

Band intensity was quantified with ImageJ software (18), for each experiment 3 different exposures were quantified and averaged.

**Immunofluorescence and Confocal Microscopy**

Samples were washed with pre-warmed PBS and fixed by incubation in pre-warmed 2% PFA in PBS solution for 10 minutes. After PBS washing, cells were permeabilized with 0.01% saponin in PBS for 5 minutes and then blocked with 5% goat serum in PBS at room temperature for 1 hour. Samples were incubated overnight at 4°C with primary antibody solution (BA8 or 3G4) used as hybridoma supernatant and diluted to 1:4 ratio in blocking buffer. On the following day, cells were washed three times in PBS, and 1-hour incubation was performed with appropriated secondary antibody solution diluted to 1:100 in blocking buffer. Samples were mounted on microscope slides with 15 µl of ProLong Gold Antifade Reagent with DAPI (Thermo fisher). Images were acquired with Nikon EclipseTi-E inverted microscope with IMA10X Argon-ion laser System by Melles Griot; all images were acquired with CFI Plan Apo VC 60X Oil (Nikon).
For p.N432D expression pattern, 14 independent experiments for a total of 2018 transfected cells were visually analyzed and assigned to one of the four categories.

For late endosome-lysosome visualization, cells were incubated for 30 minutes with 1μM LysoTracker Red DND-99 (Thermo Fisher) prior to fixation. For colocalization quantification, 12 independent experiments for a total of 248 cells were analyzed with Nikon NIS Elements software.

For ER visualization, cells were incubated for 30 minutes with 1μM ER-Tracker Green (Thermo Fisher) prior to fixation. TSHR was labeled with a red secondary antibody and staining color was then reverted to green for TSHR and red for ER with Nikon NIS Elements software in order to have graphically homogeneous data display through all the panels. For colocalization quantification, 6 independent experiments for a total of 110 cells were analyzed with Nikon NIS Elements software.

Flow cytometry

Cells were detached in Ca2+ free phosphate buffered saline (PBS), and single cell suspension was obtained by gentle pipetting. Cell suspension was transferred to FACS tubes and incubated for 30 minutes on ice in the presence of BA8 or 3G4 anti-TSHR antibody used as hybridoma supernatant and diluted to 1:4 in FACS buffer (0.1% BSA, 0.1% sodium-azide in PBS solution). Cells were washed two times in FACS buffer and labeled for 30 minutes on ice in the dark with Alexa fluor conjugated goat anti-mouse IgG (Thermo Fisher), diluted to 1:100 in FACS buffer. Cells were washed in PBS and fixed in 2% PFA-PBS. Measurements were performed with FACSCalibur flow cytometer (Becton Dickinson) on 10000 cells per sample. Six independent experiments were performed, each of them in duplicate. Data were analyzed with Flowing Software 2. For each experiment, transfected cells were gated based on the empty vector samples and mean fluorescence intensity was measured for each sample.
Functional assays

For cAMP signaling pathway activity cells were transfected with TSHR variants together with the CRE-responsive firefly luciferase construct and Renilla luciferase 40:1 mix from Cignal CRE Reporter (luc) Kit (Quiagen) following manufacturer’s instruction. Forty-eight hours after transfection cells were incubated with 0-100 U/L bovine TSH (bTSH) for 6 hours at 37°C and cAMP pathway activity was measured with Dual-Glo® Luciferase Assay System (Promega) following manufacturer’s instruction. Samples’ luminescence was measured with the Fluoroskan Ascent FL multiplate reader. Six independent experiments were performed.

Gq11/IP3 signaling pathway activity was measured with IP-One ELISA assay kit (Cisbio) following manufacturer’s instructions. Briefly, cells were incubated for 1 hour at 37°C, 5% CO2 with 100 U/L bTSH in stimulation buffer solution, followed by 30 min incubation in lysis buffer. 50 uL of each lysate were then moved to ELISA plate and reaction developed following manufacturer’s instruction. Colorimetric reaction was red at 450nm using ELx800 Absorbance Microplate Reader. Six independent experiments were performed.

Statistical analysis

All experiments were independently repeated at least three times, as indicated in the text and figure legends. After normal distribution and variance similarity evaluation, two-sided unpaired t-test (eventual Welch’s correction for groups with different variances), one-way ANOVA with Bonferroni post-hoc test, Kruskal-Wallis H test with Dunns post hoc test and Chi-square test were used as indicated in figures’ legend.

For concentration-effect curves of Gs/cAMP signaling a log(agonist) vs. normalized response - Variable slope equation was used for curve interpolation and parameters definition.
For confocal experiments, the degree of colocalization was quantified through Pearson’s correlation coefficient, as measured with Nikon NIS Elements software. Correlation was defined as strong with Pearson’s correlation coefficient bigger than 0.8, moderate when bigger than 0.5 and weak when bigger than 0.2. In all figures data are shown as mean±SEM, analyzed using GraphPad Prism 5 software and significance expressed as P values (* p < 0.05, ** p < 0.01, *** p < 0.001).
References to supplementary methods


Supplementary materials to Grassi ES et al (running title: TSHR variants degradation and functional rescue)

Supplementary Figure 1: p.N432D induces JNK phosphorylation and reduces cell viability.

A: representative images of western blotting and relative densitometric quantification showing JNK1/2 phosphorylation and total protein levels in cell transfected with different TSHR variants.

B: proliferation assays showing cells viability at 24, 48 and 72h post transfection of the different TSHR variants.

Statistical analysis: A, B n=3. Statistical significance was determined with non parametric Kruskal-Wallis H test followed by Dunn’s post hoc test.

*p<0.05, ** p<0.01 and ***p<0.001 as indicated.
Supplementary figure 2: TMAO treatments promote p.N432D homodimerization and A-subunit expression on plasma membrane.

A: representative image of TSHR western blot performed with 3G4 antibody (reblot of Fig. 4A) showing effects of TMAO treatment on high molecular weight high mannose dimers.

B: densitometric quantification of western blot experiments showing high mannose dimers, complex carbohydrates and high mannose forms of TSHR without and after TMAO treatment.

C: representative images of western blotting experiments showing the expression and maturation of the TSHR variants in Total Cell Membranes (TCM) and Plasma Membranes (PM) preparations. TSHR was stained with 28.1 antibody, VDAC was used as intracellular membranes control and E-cadherine (ECAD) was used as plasma membrane control.

D: densitometric quantification of western blot experiments showing complex carbohydrates, high mannose and A-subunit forms of TSHR without and with TMAO treatment.
Supplementary **figure 3**: glycerol treatment does not affect TSHR maturation.

Representative images of 3 independent western blotting experiment showing the expression and maturation of the TSHR variants without and after treatment with 10% glycerol. TSHR was stained with 28.1 antibody. GFP was used as transfection efficiency control and actin was used as loading control.
Supplementary figure 4: TSHR variants concentration-effect curves.

A, B: dose-response curves for Gs/cAMP signaling of TSHR WT, p.N432D and p.P449L variants upon normal conditions and after TMAO treatment. The effect is represented as percentage of TMAO treated WT maximal stimulation and expressed as Luciferase/Renilla RLU.
Supplementary table 1: in silico prediction of TSHR mutants and the respective in vitro data.

*In silico* prediction of mutation effects were obtained as described in material and methods session. TSHR variants were assigned as damaging (D) when predicted to be functionally damaging or probable functionally damaging, they were assigned as non-damaging (N) when they were predicted to be benign or neutral. *In vitro* data were obtained from extensive literature review and are reported as percentage of WT TSHR with a 5% approximation.

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