Leucine-rich repeat kinase 2 regulates autophagy through a calcium-dependent pathway involving NAADP

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Mutations in the leucine-rich repeat kinase-2 (LRRK2) gene cause late-onset Parkinson’s disease, but its physiological function has remained largely unknown. Here we report that LRRK2 activates a calcium-dependent protein kinase kinase-β (CaMKK-β)/adenosine monophosphate (AMP)-activated protein kinase (AMPK) pathway which is followed by a persistent increase in autophagosome formation. Simultaneously, LRRK2 overexpression increases the levels of the autophagy receptor p62 in a protein synthesis-dependent manner, and decreases the number of acidic lysosomes. The LRRK2-mediated effects result in increased sensitivity of cells to stressors associated with abnormal protein degradation. These effects can be mimicked by the lysosomal Ca²⁺-mobilizing messenger nicotinic acid adenine dinucleotide phosphate (NAADP) and can be reverted by an NAADP receptor antagonist or expression of dominant-negative receptor constructs. Collectively, our data indicate a molecular mechanism for LRRK2 deregulation of autophagy and reveal previously unidentified therapeutic targets.

INTRODUCTION

Autosomal-dominant mutations in leucine-rich repeat kinase 2 (LRRK2) cause late-onset familial Parkinson’s disease (PD) (1,2) which is symptomatically and neurochemically indistinguishable from sporadic PD. In addition, pathogenic mutations contribute to sporadic PD and variations increase risk for PD, suggesting that LRRK2 may be an important factor for idiopathic disease progression as well (3). The most prominent pathogenic LRRK2 mutation (G2019S), located within the kinase domain, has been consistently shown to enhance kinase activity in vitro (4–7), suggesting that it may display a dominant, gain-of-function phenotype. Overexpression of LRRK2 harbouring this disease-segregating mutation leads to neurotoxicity in vivo, and such neurotoxicity is linked to the kinase activity of LRRK2 (8–11). Thus, the pathogenic role of mutant LRRK2 seems to be dependent on catalytic activity.

Overexpression of G2019S-mutant LRRK2 in neuronal cells decreases neurite process length with a concomitant accumulation of autphagic structures (12–14), and an impaired autophagic balance is also observed upon overexpression of LRRK2 in non-neuronal and yeast cells (15,16). This is in agreement with the findings that a significant portion of endogenous LRRK2 is localized to membranous structures including ER and endosomal–lysosomal compartments (17), and implicates LRRK2 as an important regulator of macroautophagy.

Macroautophagy (hereafter referred to as autophagy) is a process whereby cytoplasmic constituents are engulfed within specialized double-membrane vesicles called autophagosomes,
and subsequently delivered to the lysosome for degradation. Autophagy normally proceeds at a low, basal rate, which plays a key role in the homeostatic clearance of old or damaged organelles and proteins (18,19). Basal autophagy is especially high in neuronal cells, and neurons undergo degeneration when basal autophagic degradation is disrupted (20,21). Autophagic failure seems to underlie a variety of neurodegenerative diseases (22), and deregulation of autophagy is evident in the brains of PD patients (23).

Despite the findings linking LRRK2 to deregulated autophagy, the exact mechanisms underlying such deregulation have not been characterized. In this work, we find that overexpression of LRRK2 causes an increase in autophagy induction through Ca$$^{2+}$$-dependent activation of a CaMKK/adenosine monophosphate (AMP)-activated protein kinase (AMPK) pathway, which can be inhibited by calcium chelation or ectopic Bcl-2. At the same time, LRRK2 causes a partial increase in lysosomal pH and a decrease of cell survival in the presence of protein aggregation-induced stress. The LRRK2-mediated effects involve activation of nicotinic acid adenine dinucleotide phosphate (NAADP)-sensitive two-pore channels (TPCs) located on acidic stores, and can be blocked by a specific antagonist. Together, our data indicate a molecular mechanism by which LRRK2 may act to regulate lysosomal Ca$$^{2+}$$ homeostasis, with downstream effects on autophagy as well as other Ca$$^{2+}$$-dependent cellular events.

RESULTS

LRRK2 overexpression causes autophagy induction

To study the effect of LRRK2 on basal autophagy, we selected a variety of constructs including full-length LRRK2 (fl-kinase) and variants bearing the pathogenic G2019S mutation (fl-G2019S) or the kinase-inactivating K1906M mutation (fl-K1906M), respectively. To evaluate whether effects are due to the kinase domain and activity of LRRK2 (12), we also used the kinase domain (kinase) or variants thereof (G2019S, K1906M) on their own, respectively. Overexpressed proteins were mostly cytosolic, wild-type (WT) and mutant proteins displayed similar overexpression levels, and no cellular toxicity was apparent 24 or 48 h upon transfection, as previously described (10) (Supplementary Material, Fig. S1).

We next addressed the effects of LRRK2 overexpression on autophagosome formation. LC3 is the mammalian homologue of the yeast Atg8 protein, which is covalently modified and re-distributes to autophagic vacuoles during induction of autophagy (23). This process can be followed by the appearance of LC3 puncta or ring-like structures and by a mobility shift from LC3I to LC3II by SDS–PAGE. Cells transiently transfected with LC3–GFP alone, or co-transfected with empty vector and analysed 48 h after transfection, display few cytoplasmic LC3–GFP punctae (Fig. 1A and B). An increase in LC3–GFP punctae was observed when co-transfecting with full-length wild-type as well as with G2019S-mutant LRRK2, but not with the inactive K1906M mutant (Fig. 1B). A similar increase in LC3–GFP punctae was observed with wild-type or G2019S-mutant, but not K1906M-mutant kinase domain only (Fig. 1A and B), indicating that the effects on autophagy are, at least in part, due to the kinase domain and activity of LRRK2. Wild-type and G2019S-mutant full-length LRRK2 increased autophagosome numbers to a similar degree and comparable with that of wild-type or G2019S-mutant kinase domain only (Fig. 1B), possibly due to saturating levels of overexpression of the various active constructs with respect to downstream targets.

As a positive control, inhibition of the mammalian target of rapamycin (mTOR) protein kinase by rapamycin was found to induce autophagy to a similar degree (Fig. 1B). The phosphatidylinositol 3-kinase inhibitor 3-methyladenine (3-MA) is thought to suppress autophagy by inhibiting the production of phosphatidylinositol-3-phosphate required for autophagosome formation (24), and was found to block the increase in autophagosome numbers mediated by rapamycin, but not by LRRK2 overexpression (Fig. 1B). Such differential 3-MA sensitivity has previously been described (13), and points towards a possible mechanistic difference between mTOR- and LRRK2-mediated autophagy induction. An increase in autophagosome numbers was further reflected by an increase in the level of endogenous LC3II (Fig. 1C), and transmission electron microscopy confirmed that the induction of LC3-positive vesicles by LRRK2 overexpression correlated with an increase in the number of autophagic structures (Fig. 1D and E).

We next used a pH-sensitive tagged LC3 construct consisting of a tandem fusion of the red, acid-insensitive mCherry (pKa < 4.5) and the acid-sensitive GFP (pKa 6.0) td-tag-LC3 (25,26). Td-tag-LC3 will emit yellow (green merged with red) fluorescence in non-acidic structures, but appear as red only in autolysosomes due to the quenching of GFP in these acidic structures (25,26). Indeed, td-tag-LC3 was found to faithfully track autophagosome maturation in transfected cells, thus allowing for the identification of possible deficits in autophagic maturation (Supplementary Material, Fig. S2). Co-transfection of td-tag-LC3 with G2019S-mutant LRRK2 kinase domain increased both early and late autophagic structures (Fig. 2A and B). The increase in both yellow and red punctae was also observed with wild-type kinase domain or with full-length LRRK2, but not with the catalytically inactive K1906M mutant (Fig. 2B). As a positive control for blocking autophagosome maturation, we used trichostatin A, an inhibitor of histone deacetylases (HDACs) known to be required for the fusion of autophagosomes with lysosomes (27). Indeed, trichostatin A treatment significantly increased the percentage of yellow punctae (and decreased the number of red punctae), in contrast to LRRK2 expression or autophagy induction by rapamycin (Fig. 2C and D). These data indicate that LRRK2 overexpression enhances autophagosome numbers and at least the initial steps of autophagosome maturation in a manner dependent on kinase activity.

LRRK2 overexpression activates the Ca$$^{2+}$/CaMKK/AMPK pathway

The increase in autophagosome numbers probably results from cytosolic events regulated by LRRK2. As recent studies imply an important role for AMPK in basal autophagy regulation (28), we evaluated the effects of LRRK2 on AMPK activity. Expression of wild-type and G2019S-mutant LRRK2 kinase domain, but not its inactive mutant, increased AMPK activity (Fig. 3A and B). This was not accompanied by inhibition of
mTORC1 activity when compared with rapamycin or torin treatment (Fig. 3C and D and Supplementary Material, Fig. S3) (29). Accordingly, inhibition of AMPK by a pharmacological inhibitor (compound C) (30) attenuated the increase in autophagosome numbers induced by LRRK2 kinase domain overexpression (Fig. 3E). Compound C also attenuated the increase in autophagosome numbers observed with ionomycin and ATP, two treatments previously shown to activate AMPK (31), but was without effect on autophagosome numbers induced by rapamycin (Fig. 3E).

Further evidence for an involvement of AMPK was gained by expressing wild-type, dominant-negative and constitutively active forms of this kinase (32). Wild-type and mutant AMPKα1 forms were overexpressed to similar degrees and without effect on cellular viability, with AMPKα2 forms expressed to a lesser degree and displaying cellular toxicity (Supplementary Material, Fig. S4). Expression of wild-type and constitutively active, but not dominant-negative, AMPKα1 induced an increase in autophagosome numbers (Fig. 4A). Importantly, the dominant-negative AMPKα1
construct was able to inhibit the increase in autophagosome numbers observed upon active LRRK2 kinase domain co-expression (Fig. 4B). Such inhibition was not observed when autophagic on-rate was increased by distinct means, indicating a specific involvement of the AMPK pathway in the LRRK2-mediated process (Supplementary Material, Fig. S4). AMPK can be activated by CaMKK-β in response to an increase in the cytosolic-free calcium $[Ca^{2+}]_c$ (31). In order to test whether LRRK2 expression regulates the Ca$^{2+}$/CaMKK-β/AMPK pathway, we first treated cells with a CaMKK-α/β inhibitor (STO-609) (33). This compound attenuated the increase in autophagosome formation induced by LRRK2 kinase domain (Fig. 4C). Such inhibition was similar to that observed when cells were treated with two stimuli which increase $[Ca^{2+}]_c$ by distinct means: ATP, which acts on P2 purinoreceptors to generate inositol 1,4,5-triphosphate (IP$_3$) that triggers the release of Ca$^{2+}$ from the ER through IP$_3$ receptor-regulated channels, or ionomycin, a Ca$^{2+}$ ionophore that serves as a mobile Ca$^{2+}$ carrier to equilibrate Ca$^{2+}$ levels across biological membranes, including plasma and ER membranes (31,34,35) (Fig. 4C). In contrast, the CaMKK-α/β inhibitor had no effect on
autophagosome numbers elicited by rapamycin treatment (Fig. 4C). Remarkably, addition of the intracellular Ca\(^{2+}\) chelator BAPTA-AM (bis-(O-aminophenoxy)-ethane-N\(_2\),N\(_2\),N\(_2\),N\(_2\)′-tetraacetic acid/tetra(acetoxymethyl)-ester) to LRRK2 kinase domain or full-length LRRK2-transfected cells for the last 2 hours completely abolished the increase in autophagosome numbers (Fig. 4D). BAPTA-AM was further able to block the ionomycin-mediated, but not the rapamycin-mediated, increase in autophagosome numbers (Supplementary Material, Fig. S4). Thus, a Ca\(^{2+}\)/CaMKK/AMPK pathway seems to be required for the increased formation of autophagosomes in response to active LRRK2 overexpression.

ER-localized Bcl-2 inhibits LRRK2-mediated autophagy induction

Bcl-2 has been reported to regulate cellular Ca\(^{2+}\) handling and Ca\(^{2+}\)/CaMKK/AMPK-induced autophagy (31,36). Thus, we tested whether expression of bcl-2 could inhibit the LRRK2-mediated increase in autophagosome numbers via its effects on Ca\(^{2+}\) homeostasis. For this purpose, we expressed wild-type bcl-2 (Bcl-WT), which is mostly mitochondrial (36,37), or mutants in which the C-terminal hydrophobic sequence has been either removed (Bcl-cyt), resulting in cytosolic expression, or exchanged to a corresponding membrane anchor from an ER-specific isoform of cytochrome b\(_5\) (Bcl-ER) or L. monocytogenes ActA (Bcl-mito), resulting in distinct subcellular localizations (31,38).

Western blot analysis confirmed equal expression levels, and only slight effects on cytotoxicity were observed when overexpressing Bcl-wt or Bcl-mito, but not Bcl-cyto or Bcl-ER, respectively (Supplementary Material, Fig. S4). Importantly, only Bcl-ER was able to fully inhibit the LRRK2 kinase domain-mediated increase in autophagosome numbers (Fig. 4E). ER-localized Bcl-2 has previously been demonstrated to lower [Ca\(^{2+}\)]\(_{\text{ER}}\) and agonist-induced Ca\(^{2+}\) leak from the ER (39). Together with the inhibitory effect of BAPTA-AM and inhibition of CaMKK on the LRRK2-mediated increase in autophagosome numbers, these results indicate that active LRRK2 may increase [Ca\(^{2+}\)]\(_{\text{ER}}\) by directly or indirectly inducing Ca\(^{2+}\) release from the ER. This release is likely mediated by IP\(_3\) receptors, as the increase in autophagosome numbers upon LRRK2 expression could not be blocked by dantrolene, a ryanodine receptor antagonist (Supplementary Material, Fig. S4).

LRRK2 alkalinizes a subpopulation of lysosomes

Overexpression of LRRK2 has previously been shown to lead to accumulation of multi-vesicular bodies and autophagosomes containing incompletely degraded material and p62 (15), a classical macroautophagy substrate (40), indicating possible defects in autophagic degradation. Indeed, overexpression of wild-type or G2019S-mutant, but not inactive K1906M-mutant, kinase domain led to a significant increase in p62 levels and p62-positive structures (Fig. 5A–C).
However, the LRRK2-mediated increase in p62 levels could be blocked by protein synthesis inhibitors (Fig. 5D), indicating that it was not reflecting changes in autophagic flux. Identical results were obtained when using ionomycin or thapsigargin (Supplementary Material, Fig. S5), suggesting that the LRRK2-mediated effects on p62 levels may be due to Ca^{2+}-dependent changes in protein synthesis (41).

A possible defect in lysosomal homeostasis, however, became apparent when employing lipid staining with BODIPY 493/503 (42), which revealed an increase in lipid droplet numbers in cells overexpressing LRRK2 kinase domain (Fig. 6A and B). In addition, lysotracker DND-99 or Lysosensor staining to determine lysosomal pH indicated a decrease in the number of positive punctae in cells overexpressing active LRRK2 kinase domain (Fig. 6C and D). This was not due to a decrease in lysosomal content (43), as no change in the levels of LAMP1 or LAMP2 were observed (Supplementary Material, Fig. S6). Interestingly, quantification of the percent colocalization of LAMP2 and lysotracker showed an increase in the number of structures positive for LAMP2, but negative for lysotracker, suggesting that LRRK2 increases pH in a subpopulation of lysosomes, without affecting the processing of lysosomal hydrolases (Fig. 6E and Supplementary Material, Fig. S6).

**LRRK2 acts through NAADP receptors**

There is considerable evidence that acidic organelles such as endosomes and lysosomes are significant stores of Ca^{2+} that can be released by the Ca^{2+} mobilizing messenger NAADP (44,45). NAADP-evoked cytosolic Ca^{2+} signals are accompanied by partial alkalinization of acidic stores (46) and amplification by ER Ca^{2+} stores (47), and lysosomal Ca^{2+} depletion causes lipid accumulation (48). That LRRK2 increases lipid droplet numbers (Fig. 6), causes an increase in lysosomal pH (Fig. 6) and regulates Ca^{2+}-dependent events in the cytosol which likely also involve the ER (Figs. 4 and 5) prompted us to consider the involvement of the NAADP pathway in mediating the effects of LRRK2.

Application of NAADP-AM was found to increase autophagosome numbers (Fig. 6F). This increase was abolished at higher NAADP concentrations, in agreement with the notion that NAADP-induced Ca^{2+} release desensitizes at high ligand concentrations (49,50). The effect of NAADP on autophagosome formation could be blocked by BAPTA-AM (Fig. 6F) and was accompanied by an increase in AMPK activity (Supplementary Material, Fig. S6). The autophagy induction observed with NAADP could be inhibited by Bcl-ER (Fig. 6G), similar to starvation-induced autophagy (36,37).
but distinct from autophagy induced by rapamycin or torin treatment (Fig. 6G). In addition, the autophagy induced by NAADP was insensitive to 3-MA, in contrast to TORC1-dependent autophagy inducers (Fig. 6H). Furthermore, NAADP caused an increase in LC3II levels (Fig. 6I), a protein synthesis-dependent increase in p62 levels (Fig. 6J), increased lipid droplet numbers (Fig. 6K) and increased lysosomal pH (Fig. 6L), thus fully mimicking the effects of LRRK2 expression.

TPCs have recently emerged as the likely targets for NAADP, with TPC1 localized to both endosomes and lysosomes, and TPC2 mainly to lysosomes (50,51). Point mutants in the putative helix of the pore-forming regions have been described, which seem to largely abolish Ca\(^{2+}\) conductance and uncover dominant-negative effects with respect to NAADP-mediated Ca\(^{2+}\) signals (51,52). Thus, we evaluated whether co-expressing wild-type or mutant TPCs may interfere with LRRK2-mediated autophagosome formation. Expression of mRFP-tagged TPC constructs indicated that wild-type and mutant versions were similarly overexpressed and localized to intracellular compartments, with TPC2 partially co-localizing with the lysosomal marker LAMP2, as previously described (51). None of the TPC constructs displayed cytotoxicity at 24 h upon overexpression, but TPC1 and its mutant were found to increase autophagosome formation (Supplementary Material, Fig. S7). Whilst consistent with the involvement of TPC1 in endolysosomal trafficking (53), this finding precluded further analysis of a possible reversal of the LRRK2-mediated increase in autophagosome numbers by TPC1 constructs. In contrast, TPC2 and its mutant did not display effects on autophagosome numbers when expressed on their own (Fig. 7A). Importantly, mutant but not wild-type TPC2 was able to block the effect of LRRK2 kinase domain on autophagosome numbers (Fig. 7A).

We next evaluated whether NED-19, a newly described NAADP antagonist (54), could reverse the LRRK2-mediated effects on autophagic flux. NED-19 on its own was without effect on autophagosome numbers, but could block the increase in autophagosome numbers mediated by active kinase domain or full-length LRRK2 (Fig. 7B). This was accompanied largely by a reversal in the LRRK2-mediated changes on AMPK activity (Supplementary Material, Fig. S7), p62 levels, LC3II levels and lysotracker-positive structures (Fig. 7C–E). Finally, when probing for an interaction between endogenous LRRK2 and GFP–TPC2, LRRK2 was found to co-immunoprecipitate with GFP–TPC2 and vice versa (Fig. 7F and Supplementary Material, Fig. S7).

Together, these data indicate that NAADP mimicks the effects of active LRRK2, whilst NED-19 application can revert those effects, and suggest that NAADP receptors may be direct targets for regulation by LRRK2.

LRRK2 sensitizes cells to cell death induced by proteasome inhibition

As the observed LRRK2-mediated effects were not cytotoxic per se, we next wondered whether they would sensitize cells towards cell death under certain conditions (55). When cells were treated with the proteasomal inhibitor MG-132, cell death was markedly accelerated in cells overexpressing active, but not inactive, LRRK2 kinase domain (Fig. 8A). The increased cell death in the presence of MG-132 in LRRK2 kinase domain-expressing cells could be blocked by NED-19 (Fig. 8B), supporting the involvement of the NAADP pathway in this process. Interestingly, application of rapamycin or torin, two specific TORC1 inhibitors, reverted the LRRK2-mediated effects on cell death in the presence of MG-132, whilst L690,440, a TORC1-independent autophagy-enhancing compound (56), was unable to do so (Fig. 8A and C). Rapamycin and torin also largely reversed the effects of LRRK2 on p62 accumulation and on the decrease in lysotracker staining (Fig. 8D and E). Measurements
in dopaminergic neuroendocrine PC12 cells yielded similar results. The NAADP-receptor antagonist NED-19 was able to revert the LRRK2-mediated increase in autophagosome numbers, the LRRK2-mediated decrease in lysosensor-positive structures and the LRRK2-mediated increase in cell death in the presence of MG-132 (Supplementary Material, Fig. S8). Together, these data indicate that additional interference with proteasomal degradation is detrimental to cells over-expressing catalytically active LRRK2, and that blocking NAADP-mediated signalling, or enhancing autophagic flux...
and/or suppressing protein synthesis or affecting lysosomal homeostasis through a TORC1-dependent pathway can overcome the enhanced susceptibility to cell death induced by LRRK2 in non-neuronal as well as neuroendocrine cells.

**DISCUSSION**

Mutations in LRRK2 are the most common genetic cause of PD, but the function of the protein remains unknown. Our study provides insight into the molecular mechanism of LRRK2 action. We have shown that LRRK2 kinase activity exerts a previously unknown regulatory role on macroautophagy that seems to involve Ca\(^{2+}\) release from lysosomes and the Ca\(^{2+}\)/CaMKK/AMPK pathway. These events seem to be mediated by activation of NAADP receptors, which causes Ca\(^{2+}\) efflux from acidic stores and is accompanied by a partial alkalinization of lysosomal store pH (Fig. 9).

In our experimental conditions, wild-type LRRK2 had a significant effect similar to pathogenic G2019S-mutant LRRK2. As mutations in LRRK2 cause PD in an autosomal-dominant manner, we overexpressed the protein to mimic a gain-of-function phenotype. It is feasible that the pathogenic mechanism of LRRK2 depends on exceeding a threshold level of activity, also supported by the finding that homo- and heterozygous carriers of LRRK2 mutations are clinically indistinguishable (3). In this context, overexpressing both wild-type or G2019S-mutant LRRK2 may reach the threshold level required for LRRK2 function and/or activity to become pathogenic. Alternatively, we may have missed a small but biologically important difference in effect, which nevertheless may have important consequences when present over many years. Finally, it remains possible that the effects described here are a wild-type function of LRRK2 unrelated to PD, even though other studies describing an impaired autophagic balance with G2019S-mutant, but not wild-type LRRK2 (12–16), are consistent with the idea of a threshold effect.

Previous studies have indicated that enhanced autophagy may mediate neurite shortening induced by G2019S LRRK2 expression, as molecular inhibition of autophagy reversed, and pharmacological activation of autophagy potentiated the effects of LRRK2 on neurite shortening (12,13). Whilst our data are consistent with the notion that LRRK2 expression causes autophagy induction, we find that pharmacological
activation of autophagy is beneficial to cell survival, contradictory to the aggravating effects of rapamycin on neurite length (13). However, the micromolar, but not nanomolar, concentrations of rapamycin employed in this study (13) leave open the possibility of secondary effects unrelated to TORC1 inhibition. In addition, even though neurite length and complexity may be relevant readouts for the cellular effects of LRRK2, enhanced stress sensitivity and cell death are equally informative for our understanding of LRRK2 function in cultured cells (55). The observed effects of LRRK2 are not sufficient to cause cell death, but increase the susceptibility of cells to further insults related to protein degradation-mediated stress, reminiscent of data obtained with LRRK2/a-synuclein double-transgenic mice (57). In addition, transgenic mice expressing G2019S-mutant LRRK2 display progressive age-dependent degeneration of dopaminergic neurons in vivo, accompanied by the accumulation of autophagic structures (14). These data imply an important regulatory role for LRRK2 in autophagic balance, which leads to eventual cell death in the presence of an additional trigger.

LRRK2 overexpression induces the formation of autophagosomes through a Ca\textsuperscript{2+}/CaMKK/AMPK pathway.

Contradictory data have emerged as to the role of cytosolic Ca\textsuperscript{2+} and autophagy, with both stimulatory (31) and inhibitory (56) effects on autophagy. Our data indicate that Ca\textsuperscript{2+} displays complex effects at different stages of the pathway, and is required in distinct intracellular locations. In this context, it is interesting to note that LRRK2 or reagents which increase intracellular Ca\textsuperscript{2+} were both found to increase p62 protein synthesis, indicating that, apart from regulation of autophagy, the LRRK2-mediated change in cytosolic Ca\textsuperscript{2+} may, at least in part, underlie some of the protein synthesis-mediated events proposed for LRRK2 (58,59).

The effects of LRRK2 on autophagy induction were found to be TORC1 independent, in agreement with recent findings that autophagy under normal nutritional conditions is TORC1 independent and regulated by AMPK and InsP\textsubscript{3}-R-mediated Ca\textsuperscript{2+} signalling from the ER (28,56,60). Furthermore, the LRRK2-mediated sensitization to cell death in the presence of proteasomal inhibition could be reversed by TORC1 inhibitors. Such reversal may be due to an increase in degradative capacity as autophagic flux is enhanced, a decrease in protein synthesis, an effect on lysosomal homeostasis (61) or a combination thereof. Indeed, evidence for all three
events were found with rapamyin. In either case, these data suggest that TORC1 inhibitors may prove beneficial in treating neurodegenerative events related to LRRK2.

LRRK2-expressing cells display a decrease in the number of lysosomes with acidic pH. Given that acidic pH is only lost in a fraction of lysosomes, further studies are warranted to elucidate whether there exists a correlation between the affected lysosomal subpopulation and the localization of endogenous NAADP receptors. In either case, the LRRK2-mediated effects on lysosomal pH are mild and do not result in a lysosomal proteolysis phenotype (62,63), and under basal conditions and in the absence of additional protein aggregation stress, these LRRK2-mediated effects do not affect cell survival.

NAADP is the most powerful endogenous Ca\(^{2+}\) mobilizing compound known to date. It seems to invoke an initial trigger release from acidic stores which is then amplified by Ca\(^{2+}\) release from ER stores, and a concomitant increase in lysosomal pH. The cytosolic increase in Ca\(^{2+}\) may result in Ca\(^{2+}\)-induced Ca\(^{2+}\) release from ER stores, and activation of CaMKK\(\beta\), followed by activation of AMPK and an increase in autophagosome numbers. These LRRK2-mediated events seem to sensitize cells to cell death in the presence of additional compounds interfering with proteasomal degradation. Detection methods and/or tools used in this study to analyse and modulate the indicated events are depicted in red, dn, dominant-negative.

**MATERIALS AND METHODS**

**Plasmid constructions**

To generate the LRRK2 kinase domain-GFPemd construct (kinase-GFPemd), the human LRRK2 kinase domain (aa 1844–2143) was PCR amplified using full-length LRRK2 as template, with EcoRI/BamHI sites on either end. VAMP2-GFPemd (64) was digested with EcoRI/BamHI and religated with the digested, PCR-amplified kinase domain cloned in-frame with the GFPemd at the C-terminal end.

To generate the LRRK2 kinase domain pCMV construct (pCMV-kinase), human LRRK2 kinase domain (amino acid 1844–2143) was PCR amplified using full-length LRRK2 as template, with SacI/BamHI sites on either end. pCMV-VAMP2 (64) was digested with SacI-BamHI and religated with the digested, PCR-amplified kinase domain.

To generate the full-length human LRRK2 pCMV vector (pCMV-fLRRK2), pGEMT-LRRK2 was digested with SacI/II to release fl-LRRK2. Quick Change Site Directed Mutagenesis was performed with the empty pCMV vector to introduce a SacI site 5’ to the fl site, followed by digestion of the vector with SacI/MluI and introduction of fLRRK2.

K1906M and G2019S mutations were introduced into all respective vectors by Quick Change Site Directed Mutagenesis, and the identity of all constructs verified by direct sequencing of the entire coding regions.

The double-myc-tagged, full-length wild-type and mutant human LRRK2 constructs were kindly provided by M. Cookson (NIH, Bethesda, USA). Full-length constructs were transformed in Sbl3 E. coli, and DNA prepared using the BACMAX DNA Purification Kit (Epicentre Biotechnologies, Madison, USA).

N-terminally myc-tagged constructs for wild-type, dominant-negative and constitutively active AMPK\(\alpha1\), as well as N-terminally myc-tagged constructs for wild-type and dominant-negative AMPK\(\alpha2\) were kindly provided by D. Carling (MRC, Imperial College London, UK). Bcl-2 and its organelle-specific mutants were kindly provided by M. Jäättelä (Institute of Cancer Biology, Danish Cancer Society, Denmark) and D.W. Andrews (McMaster University, Hamilton, Ontario, Canada). Human TPC1-mRFP, TPC1-L273P-mRFP and TPC2-mRFP have been previously described. The L265P mutation in TPC2-mRFP was generated by Quick Change Site Directed Mutagenesis and verified by direct sequencing.

The tandem-tagged LC3 construct mCherry-EF-GFP-LC3B (human) was kindly provided by T. Johansen (University of Tromso, Norway), and EGFP-LC3 (rat) was provided by T. Yoshimori (National Institute for Basic Biology, Okazaki, Japan) and J. Oliver (Institute of Parasitology and Biomedicine ‘López-Neyra’, CSIC, Granada, Spain). The G120A mutation was introduced into both tagged LC3 constructs by Quick Change Site-Directed Mutagenesis (Stratagene) and verified by direct sequencing.
Reagents

NAADP-AM and NED-19 were synthesized as previously described (54,55). Torin was a generous gift of D. Sabatini (Whitehead Institute for Biomedical Research, Cambridge, USA). 3-MA (3-methyladenine), Na3VO4, bafilomycin A1, E-64d, pepstatin A, nigericin, trichostatin A, dantrolene, puromycin and ionomycin were from Sigma. STO-609 (7-oxo-7H-benzimidazo[2,1-a]benz[de]soquinoline-3-carboxylic acid) and L690,440 ([1-(4-hydroxyphenoxo)ethylidene]bis [phosphinylidyne bis(oxyethylene)]-2,2-dimethylpropanoate) were from Tocris Bioscience (Missouri, USA). Compound C, cycloheximide and MG-132 were from Calbiochem. BAPTA-AM was from Invitrogen, rapamycin was from LC Laboratories (Woburn, USA), and chloroquine was a generous gift from L. Ruiz Pérez (Institute of Parasitology and Biomedicine ‘López-Neyra’, CSIC, Granada, Spain).

Cell culture, transfection and cell lysis

Unless otherwise indicated, experiments were performed in HEK293T cells, which were cultured as previously described (66). Where indicated, experiments were also performed in neuroendocrine PC12 cells, which were cultured and transfected as previously described (64). Cells at 70–80% confluency were transfected using Lipofectamine 2000 (Invitrogen) according to manufacturer’s specifications for 4 h, followed by addition of fresh medium. Single transfections were performed using 3.5 μg of plasmid of interest and 10 μl of Lipofectamine 2000. For double-transfections, 2.5 μg of each plasmid was used, and for triple-transfections, 1.7 μg of each plasmid was employed. The DNA ratio for co-transfections in the case of full-length LRRK2 constructs was 1:5, using a total of 10 μg of DNA.

Cells were treated with the indicated concentrations of compounds and for the indicated times, and cell extracts prepared 24 or 48 h after transfection. Cells were rinsed once in ice-cold PBS, followed by resuspension in 1 ml of lysis buffer per 100 mm dish (160 μl of lysis buffer per well of a 6-well plate) (1% SDS in PBS containing 1 mM PMSF, 1 mM Na3VO4 and 5 mM NaF). Extracts were boiled for 5 min, sonicated and centrifuged at 13 500 rpm for 10 min at 4°C. Protein concentrations of supernatants were estimated using the BCA assay (Pierce). Whenever possible, extracts were immediately resolved by SDS–PAGE without repeated freeze-thaws, as the latter was found to decrease detection of some phosphorylated proteins.

Immunofluorescence staining and confocal microscopy

Transfected cells were re-plated at a 1:3 ratio onto coverslips and processed for immunocytochemistry 24 or 48 h after transfection. Lysotracker DND-99, LysoSensor Green DND-189 and Bodipy 493/503 dyes were used essentially according to manufacturer’s instructions (Invitrogen). Fixed cells were mounted using ProLong Gold AntiFade mounting medium (Invitrogen), and images acquired on a Leica TCS-SP5 confocal microscope using a ×100 HCX PL APO CS 1.4 oil UV objective. Images were collected using single excitation for each wavelength separately [488 nm Argon Laser line and a 500–545 nm emission band pass; 543 nm HeNe Laser line and a 556–673 nm emission band pass; 405 nm UV diode and a 422–466 nm emission band pass (12.5% intensity)]. Ten to fifteen image sections of selected areas were acquired with a step size of 0.3 μm, and z-stack images analysed and processed using Leica Applied Systems (LAS AF6000) image acquisition software. The same laser intensity was used for image acquisition of individual experiments. Co-localization analysis was performed using Leica Applied Systems (LAS AF6000) image acquisition software, adjusting thresholds to 28% for each channel.

Primary antibodies included a mouse monoclonal anti-LAMP2 antibody (1:50; Santa Cruz Biotechnology), a mouse monoclonal anti-p62 antibody (1:50; BD Transduction Laboratories), a mouse monoclonal anti-myc antibody (1:10 000, Sigma) and a rabbit polyclonal anti-GFP antibody (1:1000; Abcam). Secondary antibodies included goat anti-rabbit or goat anti-mouse AlexaFluor-405-conjugated secondary antibody (1:100; Invitrogen), AlexaFluor-488-conjugated secondary antibody (1:1000; Invitrogen) or AlexaFluor-555-conjugated secondary antibody (1:1000; Invitrogen).

For detection of apoptosis, fixed cells were mounted using mounting medium containing DAPI (Vector Laboratories) and visualized on a Zeiss microscope using a ×100 oil-immersion objective. For each experiment, an average of 100 cells from random fields was quantified, and condensed or fragmented nuclei scored as apoptotic cells. Occasionally, apoptotic cells were also quantified using the DeadEnd Fluorometric TUNEL System (Promega) according to manufacturer’s specifications (Promega).

Quantitative image analysis

Autophagic activity was evaluated by quantifying the average number of GFP–LC3 punctae per cell. For this purpose, z-stacks of random fields of cells were captured at ×100 magnification on a confocal microscope. The number of GFP–LC3 punctae per cell was either obtained by manual counts performed by an observer blinded to condition, or more regularly using an NIH ImageJ macro called GFP–LC3 (13). Alternatively, random fields of transfected cells were imaged on a Zeiss microscope using a ×100 oil-immersion objective by an individual blinded to the experimental conditions, and the number of GFP–LC3 punctae per cell counted manually. Whilst the decreased resolution results in identification of overlapping or clustered punctae as a single large puncta, and thus will underestimate the number of GFP–LC3 punctae per cell, the relative changes observed using this determination were similar to those using the high-resolution analysis described above. For all determinations of LC3–GFP punctae per cell, between 30 and 50 randomly chosen transfected cells were analysed per experiment and condition. For quantification of yellow and red mCherry-EGFP–LC3 (td-tag-LC3) punctae, pictures were captured at ×100 magnification on a confocal microscope, and the number of yellow and red punctae analysed as described above from 40 randomly chosen transfected cells per experiment and condition.
**Electron microscopy**

Transfected cells were collected, resuspended in HEPES-buffered media and filtered through a 50 μm filter before FACs sorting. Sorted cells were fixed with 4% paraformaldehyde, 0.1% glutaraldehyde in 0.1 M cacodylate buffer containing 3 μM CaCl₂ and 7.5% sucrose for 15 min at room temperature. Fixed cells were washed and pelleted, then cell pellets were cut into small cubes and mixed to randomize the sample. Samples were stained with reduced osmium tetraoxide, dehydrated and embedded in resin. Sections (~80 nm) were examined with an FEI Tecnai BioTWIN electron microscope. Random cells were imaged.

To calculate relative volume densities, Photoshop software was used to combine separate images to obtain a complete view of each cell. Autophagic structures were recognized according to their morphological characteristics. A counting grid was superimposed on images at a random offset and intersections were scored according to the underlying structures, using the ImageJ plug-ins Grids and Cell Counter (NIH, Bethesda, MD, USA). The relative volume density of each category was calculated by dividing the number of intersections counted for each category by the number counted for the whole cell.

**Western blotting**

Proteins were resolved by SDS-PAGE, transferred onto polyvinylidene difluoride membranes (Hybond, GEHealthcare) and probed with primary antibodies overnight at 4°C. The following antibodies were employed: a rabbit polyclonal anti-phospho-ACC (Ser79), a rabbit monoclonal anti-phospho-AMPKα (Thr172) (40H9), a rabbit monoclonal anti-AMPKα, a rabbit polyclonal phospho-S6 (S235/236), a rabbit polyclonal phospho-p70 S6 kinase (Thr389), a rabbit polyclonal p70 S6 kinase, a rabbit monoclonal phospho-4E-BP1 (T37/46) (236B4) and a rabbit polyclonal 4E-BP1. All those antibodies were from Cell Signaling Technology, and used at a dilution of 1:500. Additional antibodies included a mouse monoclonal anti-p62 (1:500; BD Transduction Laboratories), a mouse monoclonal anti-LAMP2 (H4B4) (1:200; Santa Cruz Biotechnology), a goat polyclonal anti-cathepsin D antibody (1:100; Santa Cruz Biotechnology), a mouse monoclonal anti-cathepsin L antibody (1:100; BD Biosciences), a mouse monoclonal anti-LC3 (1:500; MBL, Japan), a rabbit polyclonal anti-LC3 (1:500; Cell Signaling Technology), a rabbit polyclonal anti-GFP (ab6556) (1:500; Abcam), a monoclonal anti-bcl antibody (ab694) (1:1000; Abcam), a monoclonal anti-tubulin antibody (clone DM1A) (1:5000; Sigma), a rabbit polyclonal anti-pan-14-3-3 antibody (K-19) (1:500; Santa Cruz Biotechnology), a rabbit monoclonal anti-LRRK2 antibody (MJF2, 1:500; Epitomics) or a mouse monoclonal C-terminal anti-LRRK2 antibody (N241A/34, NeuroMab, UCDavis, USA). Membranes were washed and incubated with secondary antibodies [anti-rabbit HRP-conjugated antibody (1:2000) or anti-mouse HRP-conjugated antibody (1:2000) (Dako Cytomation)] for 90 min at room temperature, followed by detection using ECL reagents (Roche). Unless otherwise indicated, 50 μg of cell extracts were resolved on large Protean II gels (BioRad Laboratories), whilst for phospho-p70 S6 kinase, phospho-4E-BP1, LC3 and p62, 150 μg of cell extracts were resolved on large Protean II gels (BioRad Laboratories), and westerns developed with enhanced ECL reagents (Roche). Upon chemiluminescence detection, a series of timed exposures were undertaken to ensure that densitometric analyses were performed at exposures within the linear range. Films were scanned, and QuantityOne (Biorad) was employed for densitometric analysis.

**Immunoprecipitation**

Transfected cells were washed in chilled PBS and pelleted by centrifugation. Pellets were resuspended in IP buffer (1 ml/10 cm dish) containing 1% NP-40, 50 mM Tris/HCl, pH 7.4, 150 mM NaCl, phosphatase inhibitor cocktails 2 and 3 (Sigma) and protease inhibitor cocktail (Roche) and lysates centrifuged at 13 500 rpm for 10 min at 4°C. Protein concentration of supernatants was estimated using a BCA assay, and 1 mg total protein was subjected to immunoprecipitation with either anti-GFP (5 μg, Abcam) or anti-LRRK2 (10 μg, N138/6, NeuroMab, UCDavis, USA) antibodies. For immunoprecipitations, lysates were incubated with antibodies for 1 h at 4°C, followed by addition of protein G or protein A Sepharose beads and incubation overnight. Beads were washed five times with IP buffer followed by elution with SDS sample buffer and heating at 50°C for 3 min prior to separation by SDS-PAGE.

**Statistical analysis**

Data are expressed as means ± s.e.m., and the significance of differences was assessed using unpaired t-tests. Differences were accepted as significant at the 95% level (P < 0.05).

**SUPPLEMENTARY MATERIAL**

Supplementary Material is available at HMG online.

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