Cyclophilin D Is Involved in the Regulation of Autophagy and Affects the Lifespan of *P. anserina* in Response to Mitochondrial Oxidative Stress

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The mitochondrial permeability transition pore plays a key role in programmed cell death and the induction of autophagy. Opening of the pore is regulated by the mitochondrial peptidyl prolyl-cis, trans-isomerase cyclophilin D (CYPD). Previously it was shown in the aging model organism *Podospora anserina* that PaCYPD abundance increases during aging and that PaCypD overexpressors are characterized by accelerated aging. Here, we describe a role of PaCYPD in the regulation of autophagy. We found that the accelerated aging phenotype observed in a strain overexpressing PaCypD is not metacaspase-dependent but is accompanied by an increase of general autophagy and mitophagy, the selective autophagic degradation of mitochondria. It thus is linked to what has been defined as “autophagic cell death” or “type II” programmed cell death. Moreover, we found that the previously demonstrated age-related induction of autophagy in wild-type aging depends on the presence of PaCYPD. Deletion of PaCypD leads to a decrease in autophagy in later stages of age and under paraquat-mediated oxidative stress. Finally, we report that PaCYPD is also required for mitohormesis, the beneficial effect of mild mitochondrial stress. Thus, PaCYPD plays a key role in the context-dependent regulation of pathways leading to pro-survival and pro-death effects of autophagy.

Keywords: cyclophilin D, autophagy, lifespan, aging, mitohormesis, oxidative stress

INTRODUCTION

Programmed cell death (PCD) is a cell-suicide program that is essential for proper development but is also involved in degenerative processes leading to aging and to various diseases. Due to the morphology of dying cells this process is classified into three main types: apoptotic or “type I,” autophagic or “type II,” and regulated necrotic or “type III” PCD (Schweichel and Merker, 1973; Kroemer et al., 2005). According to the Nomenclature Committee on Cell Death, extrinsic apoptosis strictly culminates in a caspase-dependent cell death, while intrinsic apoptosis leads to the release of pro-apoptotic factors from the mitochondrial intermembrane space. The latter is further subdivided into a caspase-dependent and caspase-independent cell death based on its dependency on “apoptosis-inducing factors” (AIFs) or the activation of caspases. In contrast, autophagic cell...
death is accompanied by an increased autophagic flux and can be blocked via the inhibition of autophagy. Despite its role in PCD, under physiologic conditions, autophagy acts as a pro-survival mechanism involved in metabolic adaptation and the recycling of damaged or surplus cellular components. The third type of PCD, termed regulated necrosis, is defined as a genetically controlled cell death process that lacks the classical markers of apoptosis and autophagy and often involves the activation of the kinases receptor-interacting protein 1 (RIP1) and its homolog RIP3 (Galluzzi et al., 2012).

A key process preceding the induction of autophagy and PCD is the permeability transition of the mitochondrial inner membrane resulting from an opening of the mitochondrial permeability transition pore (mPTP). Regulation of the mPTP is mediated by the mitochondrial peptidyl-prolyl cis, trans isomerase cyclophilin D (CYPD; Baines et al., 2005; Basso et al., 2005; Nakagawa et al., 2005; Schinzel et al., 2005). Oxidative stress and elevated Ca²⁺ levels lead to binding of CYPD to and opening of the mPTP. The resulting influx of water into mitochondria leads to dissipation of the mitochondrial membrane potential, swelling of the organelle, unfolding of the inner membrane, rupture of the outer membrane, release of apoptogens, and the induction of PCD via regulated apoptosis or necrosis (Schneider, 2005; Bonora et al., 2015; Izzo et al., 2016). A role of the mPTP in the degradation of mitochondrial proteins by autophagy has been reported in mammals (Carreira et al., 2010). Starvation of cardiac cells was found to result in a reduction of the mitochondrial membrane potential and the degradation of mitochondrial proteins by autophagy, which both could be prevented by treatment with the mPTP-inhibitor cyclosporine A (CsA). Furthermore, cardiomyocytes from mice overexpressing CypD exhibited an increased level of autophagy even under fed conditions.

In the fungal aging model Podospora anserina first evidence for a role of PaCYPD in aging is derived from a mitochondrial proteome analysis that revealed an age-related increase of PaCYPD (Grobe et al., 2007). Subsequent investigations showed that PaCypD deletion mutants display an increased resistance against inducers of oxidative stress and cell death. In addition, PaCypD overexpressing strains were characterized by premature nuclear condensation, cytochrome c release, reduction of the membrane potential, a severe reorganization of the mitochondrial ultrastructure, and a strong reduction of the lifespan (Brust et al., 2010a). In contrast to the role of PCD in higher eukaryotes were PCD is involved in the control of cellular homeostasis and the survival of the organism, both in the yeast Saccharomyces cerevisiae and in P. anserina apoptosis acts in the final step in the life cycle leading to death of the individual (Hamann et al., 2008). The latter is demonstrated by the finding that ablation of the two metacaspases of P. anserina and of “AIFs” lead to increased tolerance to oxidative stress and an increased lifespan. Since these data are consistent with an age-related increase of the metacaspase activity in the wild type, it was suggested that apoptosis is induced by an increase of oxidative stress in the senescent state (Hamann et al., 2007; Brust et al., 2010b). More recent data identified also a role of autophagy in aging of P. anserina (Philipp et al., 2013; Knuppertz et al., 2014) and now raise the question about mechanistic links between autophagy, CYPD, mPTP, and lifespan control.

Here we report data from a study analyzing the role of the two metacaspases and of autophagy in PaCypD overexpressing strains of P. anserina. We show that PaCypD overexpressors are characterized by an increased autophagy-dependent degradation of mitochondrial and cytosolic proteins and that this process, and not the induction of a metacaspase-dependent PCD (“type I” PCD), is responsible for the reduced lifespan of PaCypD overexpressing strains. In contrast, during physiological (“normal”) aging of the wild type, PaCYPD mediates a moderate induction of autophagy in response to oxidative stress and aging, which acts as a pro-survival pathway.

MATERIALS AND METHODS

Strains and Culture Conditions

The following P. anserina strains were used: Wild-type strain “s” (Rizet, 1953), the PaCypD-mutants ΔPaCypD, PaCypD_OEx (Brust et al., 2010a), the metacaspase deletion mutants ΔPaMca1, ΔPaMca2, ΔPaMca1/ΔPaMca2 (Hamann et al., 2007), the autophagy-deficiency mutant ΔPaAtg1 (Knuppertz et al., 2014), the PaSod3-overexpressing mutant PaSod3_OEx and the previously described mutants PaSod1::Gfp, PaSod3::Gfp (Zintel et al., 2010). These strains were used for the generation of the new mutants ΔPaCypD/PaSod3::Gfp, PaCypD_OEx/ PaSod3::Gfp, ΔPaCypD/PaSod1::Gfp, PaCypD_OEx/PaSod1::Gfp in addition to PaCypD_OEx/ΔPaAtg1 and PaCypD_OEx/ ΔPaMca1/ΔPaMca2, which were verified by Southern blot and western blot analysis (Supplementary Figures 1–8). All strains were constructed in the genetic background of the wild-type strain “s.” Strains were grown on complete medium (BMM) containing cornmeal extract (Esser, 1974) at 27°C under constant light. For spore germination, the BMM medium was supplemented with 60 mM ammonium acetate and spores were incubated at 27°C in the dark for 2 d. All used strains originated from monokaryotic ascospores isolated from irregular asci. For subsequent experiments strains were either grown on solid semi-synthetic M2 medium (M2) or in liquid complete medium (CM) at 27°C under constant light (Osiewacz et al., 2013).

Southern Blot Analysis

Total DNA was isolated according to standard protocols (Lecellier and Silar, 1994). DNA restriction, gel electrophoresis, and Southern blotting were performed according to standard protocols (Sambrook et al., 1989). As a hybridization probe specific for the phleomycin resistance gene (Blc), the 1293 bp BamHI-fragment (ER0051, Thermo Scientific) of the plasmid pKO4 (Hamann et al., 2005; Luce and Osiewacz, 2009) was used. The 727 bp NcoI/ClaI-fragment (ER0571, ER0141, Thermo Scientific) of the plasmid pKO7 (Kunstmann and Osiewacz, 2009) was used as a hybridization probe specific for the hygromycin resistance gene (Hph). For Southern blot hybridization and detection of hybridizing bands, the instructions of the supplier (Roche, Germany) were followed.
Lifespan and Growth Rate Determination

The lifespan and growth rate of monokaryotic isolates was determined using race tubes containing M2 medium as described (Kunstmann and Osiewacz, 2008). The lifespan of *P. anserina* is defined as the time period in days (d) of linear hyphal growth whereas the growth rate is defined as the measured growth in centimeters (cm) per time period in days (d). For the analysis of the growth rate and lifespan the growth front was marked every 1–3 days until death of the individuals. From these data the mean lifespan was calculated as average of all individual isolates from each strain as previously described (Osiewacz et al., 2013). To determine the lifespan under oxidative stress or in presence of cyclosporine A, petri dishes containing 30 ml M2 medium with 80 or 160 μM of paraquat (Sigma-Aldrich, 856177), CuSO₄₅₄, a combination of both or 0.05 μg/ml CsA were inoculated with mycelium from germinated spores. Plates were incubated at 27°C under constant light or in the dark in case of plates containing light sensitive cyclosporine A.

Measurement of Metacaspase Activity

For the generation of extracts of 11 days old cultures mycelium from 2 days old germinated spores of four different isolates of each strain (=biological replicates) were grown on M2 medium for 5 days at 27°C under constant light. M2 medium petri dishes covered with cellophane were inoculated with pieces of mycelium that derived from the growth front and incubated for additional 2 days. Afterwards the grown mycelia was transferred into CM liquid medium and incubated for 2 further days at 27°C under constant light and shaking at 180 rpm. The harvested mycelia were ground in liquid nitrogen and used for the isolation of total protein extracts and measurement of the arginine-specific peptidase activity using the fluorochrome-coupled peptide Z-GGR-Amc as a substrate as described (Hamann et al., 2007). Peptidase activity in total protein extracts from each strain was measured twice.

Western Blot Analysis

Cultivation of at least three different isolates of each *P. anserina* strain was carried out as described above for the determination of the metacaspase activity. For the cultivation of 6 days old strains M2 medium petri dishes covered with cellophane were directly inoculated with mycelium from germinated spores. In case of growth under oxidative stress 60 μM paraquat (Sigma-Aldrich, 856177) was added to CM liquid medium. Harvested mycelia were ground in liquid nitrogen and used for the isolation of total protein extracts with 1:100 protease inhibitor cocktail set IV (Calbiochem) as described (Osiewacz et al., 2013). One hundred micrograms total protein extracts were fractionated by 2-phase SDS-PAGE (12% separating gels) according to the standard protocol (Brust et al., 2010a). After electrophoresis, proteins were transferred to PVDF membranes (Millipore, IPFL00010). Blocking, antibody incubation, and washing steps were performed according to the Odyssey western blot analysis handbook (LI-COR Biosciences, Bad Homburg, Germany). The following primary antibodies were used: Anti-GFP (mouse, 1:10,000 dilution, Sigma-Aldrich, G6795), anti-PaCYPD (rabbit, 1:5000 dilution) previously described in Brust et al. (2010a). In all analyses, secondary antibodies conjugated with IRDye680RD (1:15,000 dilution, goat anti-mouse: LI-COR Biosciences, 926–68,070) or IRDye800CW (1:15,000 dilution, goat anti-rabbit: LI-COR Biosciences, 926–32,211) were used. The Odyssey infrared scanner (LI-COR Biosciences) was used for detection and quantification using the manufacturer’s software.

Statistical Analysis

For statistical analyses of growth rate, metacaspase activity, and quantification of the GFP protein level in western blot analyses the 2-tailed student’s *t*-test was used. For statistical analysis of lifespan the 2-tailed Wilcoxon rank-sum test was used. For all analyses statistical significance was calculated and defined as not significant (*P > 0.05); as significant (* *P < 0.05); as highly significant (** *P < 0.01); very highly significant (*** *P < 0.001).

RESULTS

Lifespan Reduction in *PaCypD* Overexpressors Does Not Rely on Metacaspase Induction

Previously it was shown that PaCYPD is involved in the induction of PCD. The analysis of a *PaCypD*-overexpressing strain with an ectopic integration of the plasmid pCypDEx1 containing *PaCypD* under the control of the constitutive promoter of the metallothionein gene *PaMtl*, identified an accelerated cytochrome c release, nuclear condensation and a decrease in membrane potential as markers for an apoptotic type of PCD (Brust et al., 2010a). However, the study did not address the dependence of the underlying processes on the two *P. anserina* metacaspases *PaMCA1* and *PaMCA2* and thus not explicitly demonstrated a metacaspase-dependent ‘type I’ PCD. In order to investigate this role, we constructed a *PaCypD_OEx/ΔPaMca1/ΔPaMca2* triple mutant in which both *PaMca* genes are deleted in the *PaCypD* overexpression background. Subsequently, we determined the arginine-specific peptidase activity of 11 days old cultures of the wild type, the *PaCypD* overexpressor, and the *PaMca1/PaMca2* double deletion in the wild-type and the *PaCypD* overexpression background (Figure 1A). The arginine-specific peptidase activity of the *ΔPaMca1/ΔPaMca2* deletion strain is very low [214 ± 27 RFU/(min*mg)] compared to the activity of the wild type [2514 ± 376 RFU/(min*mg)] indicating that the activity mainly results from the activity of *PaMCA1* and *PaMCA2*. Strikingly, the *PaCypD_OEx* mutant is characterized by a considerably lower arginine-specific activity [981 ± 359 RFU/(min*mg)] than the wild type, while the activity of the *PaCypD_OEx/ΔPaMca1/ΔPaMca2* triple mutant [258 ± 82 RFU/(min*mg)] is comparable to that of the *ΔPaMca1/ΔPaMca2* deletion strain. From these data the metacaspase activity (Figure 1B) was calculated as the difference in the arginine-specific peptidase activity between wild type and *ΔPaMca1/ΔPaMca2* strain or *PaCypD_OEx* and...

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PaCypD_OEx/ΔPaMca1/ΔPaMca2 triple mutant. Surprisingly, compared to the wild type of the same age, the metacaspase activity in 11 days old PaCypD overexpressors is strongly reduced.

Next we determined the growth rate (Figure 1C) and lifespan (Figure 1D) of the different strains on semi-synthetic M2 medium. On this medium the growth rate and mean lifespan (Supplementary Figure 9) of the wild type and the ΔPaMca1/ΔPaMca2 strain do not significantly differ from each other. In contrast, overexpression of PaCypD in the wild-type and the ΔPaMca1/ΔPaMca2 genetic background leads to a similar reduction of the growth rate and the mean lifespan. Overall these data demonstrate that the accelerated aging process of PaCypD overexpressors does not rely on metacaspase-activity and “type I” PCD.

**Lifespan Reduction in PaCypD Overexpressing Strains Depends on PaCypD-Mediated Induction of Autophagy**

Next we addressed a link of PaCypD and autophagy. For the demonstration of the vacuolar degradation of mitochondrial and cytosolic proteins via mitophagy and general autophagy, we utilized a recently established modified assay (Knuppertz et al., Frontiers in Genetics | www.frontiersin.org 4 September 2016 | Volume 7 | Article 165).
et al., 2014) that originally was developed for the yeast S. cerevisiae (Meiling-Wesse et al., 2002; Welter et al., 2010). In this assay the degradation of reporter proteins is analyzed. In our analysis, mitochondrial PaSOD3::GFP served as a marker for mitophagy and cytosolic PaSOD1::GFP for general autophagy. After autophagy-mediated degradation of parts of the fusion proteins in the vacuole, “free GFP” remains stable and can be quantified by western blot analysis. We analyzed “free GFP” in total protein extracts of two different age stages of the wild type, the PaCypD overexressor and the PaCypD deletion strain, which was obtained by homologous replacement of PaCypD by a selection marker containing a blasticidin resistance (Bsd) and phleomycin resistance (Ble) gene for selection in Escherichia coli and P. anserina (Brust et al., 2010a; Figures 2A,C). Compared to the wild type, the amount of “free GFP” derived from the PaSOD3::GFP mitophagy reporter protein was significantly lower in total protein extracts of ΔPaCypD of both 6 and 13 days old cultures. In striking contrast, compared to 6 days old cultures of the wild type, a 10-fold increase of “free GFP” was detected in extracts of PaCypD overexpressors of the same chronological age. Also, at the age of 13 days PaCypD overexpressing strains exhibit a higher amount of “free GFP” (Figure 2B). In addition, the analysis of “free GFP” resulting from degradation of the general autophagy reporter PaSOD1::GFP revealed that 6 days old wild-type and ΔPaCypD cultures display equal amounts of “free GFP”, indicating that the basal level of general autophagy is independent of PaCYPD. In striking contrast, we verified the age-related increase in general autophagy firstly reported for the wild type of P. anserina (Knuppertz et al., 2014) and found that it is PaCYPD-dependent. In ΔPaCypD strains “free GFP” of 11 days old cultures is significantly lower than in wild-type strains of the same age and no significant difference of “free GFP” is observed in 6 and 11 days old ΔPaCypD cultures. In addition, compared to 6 days old wild-type cultures, PaCypD overexpressors display a 3.5-fold increase of “free GFP” in 6 days cultures and a 4.6-fold increase at an age of 11 days (Figure 2D). The mitophagy reporter PaSOD3::GFP may also be degraded (together with cytosolic compounds) unspecifically to some extent in the PaCypD overexpressor. However, since the increase in degradation of PaSOD3::GFP is much higher than the increase in PaSOD1::GFP degradation (10-fold vs. 3.5-fold) a preferential degradation of PaSOD3::GFP via mitophagy can be concluded.

Next we investigated the impact of autophagy on growth and lifespan of PaCypD overexpressors. Ablation of PaATG1, an essential component of the autophagic machinery controlling early steps in autophagosome formation, impairs the pro-survival function of autophagy and results in a reduction of the growth rate and a short-lived phenotype (Knuppertz et al., 2014). Most strikingly, we found that growth rate and lifespan of the PaCypD_OEx/ΔPaAtg1 double mutant is not further reduced in comparison to the two single mutants (Figures 2E,F, Supplementary Figure 10), indicating an epistatic relation of PaAtg1 and PaCypD. Thus, a pro-survival role of autophagy, as it is found in the wild type, can be excluded in the PaCypD overexpressor. We conclude that lifespan reduction in this mutant is caused by the CYPD-dependent strong induction of general autophagy and mitophagy. PaCYPD thus appears to be involved in the switch from a pro-survival to a pro-death role of autophagy. Moreover, as indicated by the lifespan reduction in the PaCypD_OEx/ΔPaAtg1 double mutant, a complete suppression of autophagy in the PaCypD overexpression background is also not beneficial clearly demonstrating that balancing of autophagy flux is important for cellular homeostasis.

PaCYPD Is Involved in Paraquat-Mediated ROS Signaling

After the identification of a role of PaCYPD in the induction of ROS signaling, we set out to investigate the underlying mechanisms in more detail. Since a role of ROS in aging and in the regulation of CYPD-induced mPTP is well documented (Linard et al., 2009; Davalli et al., 2016), we next analyzed the impact of oxidative stress on the growth rate of the wild type and different PaCypD mutants (Figure 3A). We induced global oxidative stress in all cellular compartments via the addition of CuSO₄ to the growth medium. In addition, the herbicide paraquat was used to induce mitochondrial stress. Under standard conditions without the application of the two stressors, the growth rate of ΔPaCypD and PaCypD_OEx strains is slightly reduced compared to the growth rate of the wild type. On medium containing 80 and 160 µM CuSO₄, respectively, the growth rate of ΔPaCypD and the wild type do not differ. Growth of PaCypD_OEx is only slightly decreased compared to the wild type, indicating that the tolerance to Cu²⁺-induced global cellular oxidative stress is independent from PaCYPD. In contrast, compared to the wild type, 80 and 160 µM paraquat in the growth medium resulted in a significant increase of the growth rate of ΔPaCypD and a decreased growth of PaCypD_OEx on medium containing 80 µM paraquat. It appears, that ΔPaCypD strains are more tolerant to paraquat-mediated mitochondrial ROS stress. While all three investigated strains were able to grow on 160 µM CuSO₄ or 160 µM paraquat, respectively, only the wild type and the PaCypD deletion strain were tolerant to the addition of 80 µM of both ROS stressors. Under these conditions, growth of PaCypD_OEx was almost completely inhibited. These data are consistent with observations in higher eukaryotes in which CYPD facilitates ROS-induced mPTP-opening leading to the induction of cell death.

To validate this latter role in P. anserina, we tested whether the reduced lifespan of a mutant overexpressing PaSod3, encoding the mitochondrial superoxide dismutase, can be restored by the CYPD-inhibitor cyclosporine A. The mutant is characterized by increased ROS stress (Grimm and Osiewacz, 2015). Upon application of 0.05 µg/ml cyclosporine A to the growth medium, the growth rates of the wild type and PaSod3_OEx are significantly decreased, but do not differ from each other (Figure 3B). The cyclosporine A-mediated impairment of the growth rate is consistent with earlier findings reported for Neurospora crassa (Tropschug et al., 1989; Bardiya and Shiu, 2007) and was assumed to result from a PaCYPD/CsA interaction product, since the growth rate of
FIGURE 2 | Autophagy-dependent degradation of mitochondrial and cytosolic marker proteins in wild-type strain and different PaCyD mutants during aging. (A) Monitoring mitophagy by western blot analysis using the mitochondrial protein PaSOD3::GFP. 6 and 13 d old wild-type (WT), ΔPaCyD and PaCyD_OEx (Continued)
strains expressing PaSod3::Gfp were cultured for 2 d in CM medium. “Free GFP” (indicated by arrow) and PaCYPD were monitored by immunoblotting with anti-GFP and anti-PaCYPD in 100 µg total protein extracts. The positions of molecular mass markers are indicated on the right. (B) The GFP protein levels of three different isolates of each strain were quantified and normalized to the Coomassie-stained gels (loading control). The protein amount present in the 6 d old wild type was set to one. (C) Monitoring autophagy as described in (A) except using 6 and 11 d old strains expressing PaSod1::Gfp. (D) Quantification of “free GFP” derived from the cytosolic marker PaSOD1::GFP as described in (B). (E) Growth rate and (F) lifespan of wild type (n = 20), PaCypD_OEx (n = 20), ∆PaAtg1 (n = 16), and the double mutant PaCypD_OEx/∆PaAtg1 (n = 15) on M2 medium. Data represent average ± SEM (2-tailed student’s t-test), *P < 0.05, **P < 0.01, ***P < 0.001.

**FIGURE 3 |** (A) The growth rate of 2 d old wild-type, ∆PaCypD and PaCypD_OEx strains (n = 10) was monitored for 13 d on M2 medium in the presence of the ROS stressors paraquat (PQ), CuSO₄, both or without ROS stressors (control: C). (B) Growth rate and (C) lifespan of wild type (n = 33) and PaSod3_OEx (n = 20) on M2 medium with 0 or 0.05 µg/ml cyclosporine A. Data represent average ± SEM (2-tailed student’s t-test). Not significant (n.s.), **P < 0.01, ***P < 0.001.

Under Physiological Conditions
ROS-Mediated Autophagy Is PaCYPD-Dependent and Acts as a Pro-Survival Pathway

Since CYPD-mediated opening of the mPTP can be induced by ROS and since we recently found that autophagy is also induced by ROS in *P. anserina* (Knuppertz et al., submitted), we next analyzed whether the induction of autophagy depends on PaCYPD. In this study we compared “free GFP” levels in 6

ΔPaCypD is unaffected by CsA (Brust et al., 2010a). Moreover, chronic treatment with cyclosporine A does not affect the mean lifespan of the wild type. In striking contrast, the lifespan of PaSod3_OEx is significantly increased from 16.3 ± 2 to 20.4 ± 3 days (Figure 3C, Supplementary Figure 11). Together these data suggest that PaCYPD-inhibition or deficiency mediates an increased tolerance to mitochondrial but not to global oxidative stress. In addition, premature death caused by high mitochondrial oxidative stress can be counteracted by the CYPD-inhibitor cyclosporine A.
days old cultures of wild-type and ∆PaCypD strains challenged by 60 µM paraquat (Figure 4A). As a control we used the strains of the same age grown in medium without paraquat from the age-related study described above (Figure 2D). In the wild type, paraquat led to a two-fold increase of “free GFP” (Figure 4B). Although paraquat also led to increased “free GFP” in ∆PaCypD, the level is significantly lower than in the wild type indicating that the paraquat-mediated induction of autophagy is indeed, at least partially, dependent on PaCypD.

Next we investigated whether the beneficial autophagy-dependent effect of mild paraquat stress identified in a parallel study (Knuppertz et al., submitted) also depends on PaCYPD. We
compared the lifespans of the wild type and the \( \Delta PaCypD \) mutant grown on standard growth medium containing increasing amounts of paraquat (Figure 4C). Strikingly, we found that the lifespan-extending effect of paraquat observed for the wild type is strongly reduced in the \( \Delta PaCypD \) strain. On medium containing 20 \( \mu \)M paraquat the difference in mean lifespan is significant (Figure 4D). In contrast, at 80 \( \mu \)M the difference in mean lifespan is not significant and under harsh paraquat stress (i.e., 150 \( \mu \)M) the lifespan is strongly reduced in both strains. These results are consistent with previous investigations in other organisms like \textit{S. cerevisiae} or \textit{Caenorhabditis elegans} (Cypser and Johnson, 2002; Mesquita et al., 2010; Yang and Hekimi, 2010) and provide mechanistic clues explaining the beneficial effect of mild stress via the induction of adaptive responses leading to increased stress tolerance. They support a beneficial role of \textit{PaCYPD} in the mediation of mitohormesis.

**DISCUSSION**

Previous work identified PCD as the final step in the lifetime of the fungal aging model \textit{P. anserina}. Different components of a PCD machinery including apoptosis inducing factors (AIFs), poly(ADP-ribose) polymerase (PARP), metacaspases (MCAs), and the mitochondrial peptidyl prolyl-cis, trans-isomerase cyclophilin D (CYPD) were found to be involved in PCD and impact \textit{P. anserina} lifespan (Grobe et al., 2007; Hamann et al., 2007; Brust et al., 2010a,b; Müller-Ohldach et al., 2011; Daum et al., 2013; Strobel and Osiewacz, 2013). In the current study, we report a role of one of these components, \textit{PaCYPD}, in the induction of autophagy, a pathway that previously was identified as a pro-survival mechanism involved in longevity assurance in \textit{P. anserina} (Philipp et al., 2013; Knuppertz et al., 2014).

We started our study with the investigation of a short-lived mutant in which \textit{PaCypD} is strongly overexpressed. The mutant was previously shown to be severely impaired in mitochondrial ultrastructure, membrane potential and other characteristics of induced PCD (Brust et al., 2010a). We now demonstrate that this mutant accelerated aging and PCD is metacaspase-independent and is characterized by a strong induction of mitophagy and general autophagy. In addition, we observed that the strong increase in autophagic flux does not function as a pro-survival mechanism but is rather the cause of the deleterious effects of \textit{PaCypD} overexpression. Death in this mutant thus confers to what has previously been defined as “type II” PCD or “autophagic cell death” and general autophagy and mitophagy represent pro-death pathways. Our data about a role of \textit{PaCYPD} and the mPTP in the induction of autophagy are well consistent with what has been previously reported in mammals. For instance, in rat hepatocytes serum deprivation and glucagon caused the depolarization of mitochondria, followed by the sequestration of these mitochondria by autophagosomes, a process that can be prevented by the CYPD-inhibitor CsA (Elmore et al., 2001; Rodriguez-Enriquez et al., 2006). In addition, it was shown in myocytes from mice overexpressing \textit{CypD} that autophagy is enhanced even under fed (non-starvation) conditions (Carreira et al., 2010). The pro-death role of \textit{PaCYPD}-mediated mitochondrial permeability transition under severe stress is further emphasized by the lifespan-extending effect of the CYPD-inhibitor CsA in a \textit{P. anserina} mutant overexpressing \textit{PaSOD3}, which suffers from high oxidative stress and is also short-lived (Zintel et al., 2010; Grimm and Osiewacz, 2015). Although it is currently not explicitly shown whether death of this mutant is mediated by an autophagic or apoptotic type of PCD, the characteristics of the mutant suggest that it is likely that \textit{PaCYPD}-mediated “type II” PCD is executed in this mutant.

Apart from the role of \textit{PaCYPD} and autophagy in situations of severe stress our current study also casts new light on the role and regulation of autophagy during physiologic (“normal”) aging of the wild type and in situations of mild oxidative stress. In particular, we found that the age-related increase in general autophagy first reported for the wild type of \textit{P. anserina} (Knuppertz et al., 2014) is impaired in \textit{PaCypD}-deficient strains. Moreover, compared to the wild type, \( \Delta PaCypD \) strains exhibit a reduction of general autophagy in response to paraquat-mediated oxidative stress. Since superoxide is known to be required for starvation-induced autophagy (Chen et al., 2009) these results are well consistent with the observation that starvation of mice failed to induce autophagy in \textit{CypD}-deficient cardiomyocytes (Carreira et al., 2010). Accordingly, in a \textit{P. anserina} mutant lacking the mitochondrial superoxide dismutase \textit{PaSOD3}, mitophagy is strongly induced. This induction was concluded to result from the increased mitochondrial superoxide load since treatment with paraquat was shown to induce both autophagy and mitophagy (Knuppertz et al., submitted). In \textit{P. anserina} we obtained first evidence about a role of \textit{PaCYPD} in pro-survival signaling as a response to mild oxidative stress and its impact on longevity. Although such a hormetic response is known to be triggered by many compounds like rapamycin, spermidine, and resveratrol (Rubinsztein et al., 2011) which are involved in autophagy induction, it is important to consider that hormesis has a pleiotropic basis and is not restricted to autophagy. Moreover, the role of \textit{PaCYPD} in hormesis is very complex since the physiological role of \textit{PaCYPD} has been implicated in the regulation of \textit{Ca\textsuperscript{2+}}-homeostasis and mitochondrial metabolism (Elrod and Molkentin, 2013).

In mammals, a role of \textit{CYPD} in pro-survival and pro-death pathways has been implicated in tissue damage resulting from ischemia-reperfusion injury (IRI), the damage caused by reperfusion of tissue following a period of restricted blood supply (ischemia). Induction of PCD mediated by opening of the mPTP (high-conductance state) is a critical determinant contributing to IRI and can be counteracted by treatment with CsA or genetic ablation of \textit{CYPD} (Baines et al., 2005). In addition, one of the most protective mechanisms against IRI is ischemic preconditioning (IPC). IPC is pro-survival signaling elicited by transient, reciprocal episodes of ischemia/reperfusion prior to a major ischemic event and represents a special form of hormesis (Zhao et al., 2013). It was shown that IPC is dependent on transient mPTP-openings (low-conductance state; Hausenloy et al., 2010) and is defective in \textit{CypD}-deficient mice (Lim et al.,...
2007). Interestingly, the ability to undergo IPC decreases in mice with age while treatment of IRI with CsA remains effective (Peart et al., 2014). Thus, the dual role of PaCYPD in the aging process of *P. anserina* exhibits many similarities to the roles of CYPD in mammalian systems.

Overall, in the current study, we integrated PaCYPD and mPT into the complex molecular network involved in cellular quality and lifespan control. It appears that under “normal” conditions autophagy, which is regulated by ROS and the contribution of PaCYPD-mediated mPT, increases during aging as a pro-survival response counteracting age-related accumulation of impairments. This process can experimentally be induced via challenging “healthy” strains by mild stress and leads to the beneficial hormetric effect. The adaptive capacity of this pro-survival system is however restricted. Passing threshold levels it leads to the activation of components of “type I” PCD (e.g., metacaspases) and to death of senescent cultures. Under special conditions of excessive, non-physiological stress the system can turn to a system in which no attempts to counteract molecular impairments are induced anymore (i.e., induction of beneficial levels of autophagy). Under these conditions it leads to “type II” cell death and a rapid deterioration.

**AUTHOR CONTRIBUTIONS**

HO and PK designed the study, PK, AJ, and AH performed experiments and analyzed the data. HO and PK wrote the manuscript. HO supervised the study. All authors read the final version of the manuscript.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fgene.2016.00165

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