Dietary fat composition influences glomerular and proximal convoluted tubule cell structure and autophagic processes in kidneys from calorie-restricted mice

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Summary
Calorie restriction (CR) has been repeatedly shown to prevent cancer, diabetes, hypertension, and other age-related diseases in a wide range of animals, including non-human primates and humans. In rodents, CR also increases lifespan and is a powerful tool for studying the aging process. Recently, it has been reported in mice that dietary fat plays an important role in determining lifespan extension with 40% CR. In these conditions, animals fed lard as dietary fat showed an increased longevity compared with mice fed soybean or fish oils. In this paper, we study the effect of these dietary fats on structural and physiological parameters of kidney from mice maintained on 40% CR for 6 and 18 months. Analyses were performed using quantitative electron microscopy techniques and protein expression in Western blots. CR mitigated most of the analyzed age-related parameters in kidney, such as glomerular basement membrane thickness, mitochondrial mass in convoluted proximal tubules and autophagic markers in renal homogenates. The lard group showed improved preservation of several renal structures with aging when compared to the other CR diet groups. These results indicate that dietary fat modulates renal structure and function in CR mice and plays an essential role in the determination of health span in rodents.

Key words: aging; calorie restriction; dietary fat; kidney; mice.

Introduction
Aging can be defined as a time-dependent degenerative process caused by accumulated damage that leads to cell dysfunction, tissue failure, and death (Campisi, 2013). Although the action of free radicals (many of them produced at the mitochondria) may contribute to aging, the mechanisms through which this occurs are still not entirely known. Recently, several hallmarks have been proposed to explain the molecular and physiological basis of aging and mitochondrial dysfunction seems to play a central role in this process (López-Otín et al., 2013; González-Freire et al., 2015).

To analyze the physiological basis of aging, several experimental procedures have been developed. Among them, the reduction of calorie intake without malnutrition, also known as calorie restriction (CR), has been shown to be the most robust nongenetic or pharmacological approach to study this phenomenon (Weindruch & Walford, 1988). A reduction in calorie intake (typically 20-40% of the ad libitum fed controls) has been reported to increase lifespan and to prevent cancer, diabetes, hypertension, and other age-related diseases in a wide range of animals, including non-human primates and humans (Colman et al., 2009; Mattison et al., 2012). Although the mechanisms by which CR operates are not completely understood, it is often assumed that the anti-aging action of CR is partially based on its ability to suppress oxidative stress and maintain the cellular redox status to provide optimal cell signaling processes and normal gene expression (Chung et al., 2013). Also, CR has been proposed to induce biogenesis of efficient mitochondria (López-Lluch et al., 2008).

We have recently confirmed that the composition of dietary fat modulates longevity of mice fed CR diets (López-Domínguez et al., 2015a). Animals fed a 40% CR diet with lard (CLR, high in saturated and monounsaturated fatty acids) as the primary dietary fat had extended lifespan compared to CR animals consuming diets with either soybean oil (CRS, high in n-6 polyunsaturated fatty acids, PUFAs) or fish oil (CRF, high in n-3 PUFAs) as the primary lipid sources (López-Domínguez et al., 2015a). However, the influence of dietary fat composition on physiological function and health span are not known.

According to the ‘Mitochondrial Free Radical Theory of Aging’ (Miquel et al., 1980), accumulation of reactive oxygen species (ROS) in mitochondria (the subcellular organelle with the highest rate of ROS production) results in damage not only to mitochondrial DNA and proteins, but also to membrane phospholipids, and this oxidative damage is decreased in animals maintained on CR (Youngman et al., 1992; Pamplona et al., 2002). An inverse correlation between lifespan and the degree of membrane phospholipid unsaturation has been proposed (Pamplona et al., 2002; Hulbert, 2003), with PUFAs being more susceptible to peroxidation and other modifications which result in the accumulation of oxidative injury in membranes containing these fatty acids. The decreased content of long-chain PUFAs in mitochondria isolated from different organs after CR seems to support this idea (Yu et al., 2002). According to these results, we hypothesized that the extended longevity found in the CRL-fed group correlates with the higher MUFA content in mitochondria from these animals as detected in hepatocytes and skeletal muscle (López-Domínguez et al., 2013, 2015b). Furthermore, the decreased susceptibility of membranes to phospholipid peroxidation may improve mitochondrial function, a phenomenon likely linked to lifespan extension (Jové et al., 2014).
Kidney has been considered as an essential organ to understand the aging process, and renal function has been suggested to be one of the major predictors of longevity (Hediger, 2002). Thus, as renal function declines with age, several structural and functional changes have been reported to occur in renal glomeruli and in proximal convoluted tubule (PCT) epithelial cells (see, for example, Martin & Sheaff, 2007; Wiggins, 2012; Bolignano et al., 2014). However, relatively little is known about the influence of dietary fats on changes in renal structure and function in CR animals.

Besides the potential role of CR in mitigating oxidative stress, autophagy is also a crucial phenomenon to explain CR effects on longevity and healthy aging (Rajewat et al., 2009; Speakman & Mitchell, 2011 and Madeo et al., 2015). By this mechanism, different aged subcellular structures which accumulate molecular damage are degraded through a lysosomal pathway and the resulting products are released into the cytosol for recycling or to supply energy during starvation periods (Cuervo, 2004). Deregulation of autophagy has been shown to be involved in the pathogenesis of a number of renal disorders, many of them directly related to aging (Huber et al., 2012). Kidney aging markers were delayed or even reversed by CR (Wiggins et al., 2005; McKiernan et al., 2007), but little is known about the impact of the different dietary constituents on kidney aging in CR mice. In this study, we analyze structural and ultrastructural changes in kidney mediated by different dietary fats in CR mice. Moreover, in an attempt to establish possible links between healthy renal aging and health span expansion mediated by dietary fat under CR conditions, we have also analyzed changes in the expression of proteins related to mitochondrial biogenesis and autophagy processes.

Results

Physiological parameters and p16 expression levels

Body weight and urea and creatinine serum levels after 6 and 18 months of dietary intervention are shown in Table 1. As expected, mice sequentially lost weight in the CRS compared to the control (CON) group and there were no differences between CR groups at either 6 or 18 months (Table 1). A similar result was found for serum urea levels, which were decreased in the CRS group at both time periods. Under CR, dietary fat did not induce changes in this parameter. On the other hand, creatinine levels increased in CRL after 6 months of intervention but decreased to reach similar values to the other CR groups at 18 months. As occurred for urea, dietary fat did not induce significant changes among CR groups at 18 months (Table 1).

The expression level of p16, a marker of tissue aging, changed during CR. Six months of CR induced a decrease in this parameter when compared with the CON group, but no differences were found between CRS and control (CON) mice at 18 months (Fig. 1A). When comparing the different CR groups, a significant increase in p16 expression level was found at 18 versus 6 months regardless of the dietary fat source (see Fig. 1B) but no changes were detected in 6 or 18 months when comparing among the different CR groups (Fig. 1B).

Renal corpuscle and glomerular filtration barrier morphology

Examination of semi-thick sections of renal cortex from 6-month-old CON mice allowed us to visualize renal corpuscles showing a typical morphology. When comparing corpuscle structure in CON group with that obtained from CRS mice after 6 months of CR, no changes were detected since virtually all the glomeruli showed an unaltered morphology (Fig. 2A,B). However, in 18-month-old CON and CRS animals, sections showed a high number of glomerulosclerosis figures (about 50–55%) with no appreciable differences in percentage among these two dietary groups (Fig. 2C,D). However, an ultrastructural analysis performed on glomeruli with unaltered morphology showed striking differences between CON and CRS mice.

Glomerular basement membrane (GBM) thickness was measured on high-magnification electron micrographs (see Fig. 2E,F), and the results are depicted in Fig. 2G. Aging resulted in a significant increase (P < 0.001) of this parameter in all of the diet groups. Differences were also found when comparing control and CRS diets at 6 and 18 months of CR. After 6 months, dietary fat also induced changes in GBM thickness among the CR groups. Thus, a significant increase was found in CRS- and CRF-fed animals in comparison with CRL, which showed the lowest value. When comparing animals at 18 months of CR, we found that basal membrane enlargement due to aging was reduced to different degrees depending on the dietary fat, with CRL mice showing the thinnest GBM in their glomeruli compared to the others CR groups (see Fig. 2G).

Glomerular filtration slits (FS), which can be observed as narrow spaces between podocyte processes (see Fig. 2E,F), were also measured, and the results are displayed in Fig. 2H. In CON mice, aging induced a significant decrease in this parameter that was not observed in CRS group where FS width increased with age (Fig. 2H). Similarly, all CR groups showed a decrease in FS width with aging. Of note, the decrease observed in the CRL groups was not as prominent as occurred in the CRF group. In 6-month-old mice, FS showed similar widths regardless of diet except for CRS in which mean slit width was significantly decreased

### Table 1

|                     | CON       | CRL       | CRS       | CRF       |
|---------------------|-----------|-----------|-----------|-----------|
| Weight (g)          | 37.70 ± 1.33a |
|                     | 32.56 ± 2.58 |
|                     | 37.71 ± 0.78 |
|                     | 29.19 ± 0.75 |
| Serum urea (mg dL⁻¹) | 36.73 ± 1.43c |
|                     | 36.24 ± 2.14d |
|                     | 31.73 ± 1.19 |
|                     | 32.96 ± 3.87 |
|                     | 31.8 ± 1.85  |
|                     | 29.32 ± 1.82 |
| Serum creatinine (mg dL⁻¹) | 0.68 ± 0.06   |
|                     | 0.61 ± 0.12  |
|                     | 0.85 ± 0.08d |
|                     | 0.54 ± 0.07  |
|                     | 0.61 ± 0.07  |
|                     | 0.61 ± 0.06  |
|                     | 0.50 ± 0.09  |
|                     | 0.63 ± 0.07  |

*P < 0.05 vs CON 18 months and P < 0.01 vs CRS 6 months.

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GBM thickness and FS width was found both in 6- and 18-month interventions, and, interestingly, a similar result was obtained when comparing FS with PFP. However, PFP width and GBM thickness positively correlated in all of the dietary groups. These correlations seem to be independent of the dietary fat but clearly depend on the animal age (see Fig. S4A,B,C).

Mitochondrial mass and ultrastructure in PCT cells

The examination of thin sections of renal cortex from 6- and 18-month-old control and CR mice revealed among other structures a relatively high number of glomerular corpuscles as well as cross-sectioned proximal convoluted tubules (PCT) displaying a typical structure with epithelial cells showing a well-developed apical brush border, basal or central nuclei and a high number of mitochondria profiles. In Fig. S1 (Supporting information), we show one of these sections as an example of the materials used in this work. First, we performed a planimetric analysis of PCT epithelial cells and nuclei from different diet groups. Epithelial cellular size decreased during aging in control mice but remained unaltered in the CRS group (Fig. 3A). However, when comparing cellular area among the different CR groups, CRF showed cell sizes significantly higher than those of CRL, which decreased during aging, and CRS at both 6 and 18 months (Fig. 3A). Neither aging nor diet affected nuclear size when comparing 6- versus 18-month-old mice in any diet group (Fig. 3B). Nevertheless, in 6-month-old animals some differences appeared when comparing the CR groups with CRS having the smallest and CRF the largest nuclei (Fig. 3B).

In PCT epithelial cells, mitochondria appear as numerous electron-dense structures spread throughout the cytoplasm regardless of mouse age or dietary group. Some examples of mitochondrial appearance in PCT cells are included in Fig. S2A–D (Supporting information). After planimetric and stereological analyses of mitochondria, striking differences were found in the experimental groups. Mean mitochondrial volume increased during aging in CON mice (Fig 3C), and the same effect was observed in the CRS group. CRL and CRF followed a nearly identical pattern to the CRS group. Similar changes were observed when comparing mitochondrial area and major and minor diameters from the 3 CR groups (not shown). Aging also induced a significant decrease in mitochondrial circularity coefficient in CON and CRS mice (Fig. 3D). In addition, significant differences were also found when comparing the CR groups of animals fed with the different dietary fats. In this case, circularity coefficient was higher in CRF compared to all the other CR groups (Fig 3D).

The stereological parameters volume density (Vv; i.e., cell volume fraction occupied by mitochondria) and numerical density (Nv; i.e., number of mitochondria per cell volume unit) also changed during aging and/or CR. In CON animals, aging resulted in a decrease of Vv, a phenomenon that was not observed in CRS mice (Fig. 3E). However, when comparing the different CR groups, a significant decrease of Vv was found in the CRL group while CRF remained unaltered compared to CRS (see Fig. 3E). Conversely, aging affected Nv in different ways depending on the experimental group. Thus, in CON mice Nv did not change significantly in 6- versus 18-month-old animals, but markedly decreased in CRS mice (Fig. 3F). In this group, we also found increased Nv after 6 months when compared to CON. When comparing Nv among the CR dietary groups at a given age, we found the highest values in CRS after 6 months of dietary intervention and in CRL after 18 months. In aged animals, a decreasing linear trend CRL > CRS > CRF was found for Nv (see Fig. 3F). Of note, statistical analyses showed a clear correlation between GBM thickness and mitochondrial Vv and Nv.
in PCT cells in such a way that a thicker GBM corresponded with lesser mitochondrial mass in PCT cells. A similar correlation was observed when comparing glomerular FS width and mitochondrial Vv in PCT cells (see Fig. S4D,E,F).

The observation of mitochondrial size and mass variations among the different ages and diets led us to explore possible changes in mitochondrial dynamics and/or biogenesis, and thus, we performed an expression analysis of PGC-1α, the regulatory master of these processes, and its downstream transcription factors NRF1 (nuclear respiratory factor 1) and TFAM (mitochondrial transcription factor A). Our results show that protein expression of all three proteins did not change significantly at 6 versus 18 months in control mice but were markedly decreased with aging in CRS mice (Fig 4A,C,E). After 6 months of CR, PGC-1α, and NRF1 remained unaltered in all the three CR groups but TFAM increased significantly in CRS compared with CRL (Fig 4B,D,F). However, after 18 months of CR, the expression levels of PGC-1α, TFAM and NRF1 were higher in CRL (P < 0.01) compared to CRS and CRF (see Fig 4B,D,F).

**Autophagy ultrastructural observations**

Electron microscopy images showing typical structures of autophagy (see, for example, Hartleben et al., 2010 and Kume et al., 2010) were found in podocytes (Fig. 5A,B) and in PCT epithelial cells (Fig. 5C–G). These figures consisted of cytoplasmic portions surrounded by membranes showing irregular shape and content which frequently appeared as typical myelinlike figures. Although these structures appeared in all of the experimental groups regardless of age or feeding condition, they were more abundant in 18-month-old animals (Fig. 5A–H). In some cases, PCT showed considerable sized myelinlike structures occupying most of the cellular space (see Fig 5C). A quantitative analysis of these
Changes in expression levels of LC3 were also evaluated. The pre-LC3 form is cleaved into the cytosolic form LC3-I, which is then conjugated to phosphatidylethanolamine to form LC3-II (see, for example, Cui et al., 2012, 2013). The ratio LC3-II/LC3-I + LC3-II is correlated with autophagic flux. LC3 ratio remained unchanged in CON or CRS animals during aging. However, a significant increase LC3 ratio was observed in the CRS compared to CON group (Fig. 6C). When comparing the effects of the dietary fats, we found similar levels of LC3 ratio at 6 and 18 months in the CLR and/or CRS groups. However, CRF animals showed increased ratios at 6 and 18 months of CR when compared to the corresponding CRS groups (Fig. 6D).

**Discussion**

We have recently shown that 40% CR extends lifespan in mice. However, differences in longevity were found among CR groups depending on the source of dietary fat. Thus, lard extended longevity compared to soy and fish oils (López-Domínguez et al., 2015a) pointing to a role of specific dietary components in determining lifespan of mice fed CR diets. To assess the precise effects of this nutritional intervention at a tissue level, we analyzed the impact of dietary fat in different organs and tissues from animals fed these same diets (see Khraiwesh et al., 2013 and Khraiwesh et al., 2014 and López-Domínguez et al., 2013, 2014).
In this paper, we studied kidney structure and biology in an attempt to determine the possible role this organ plays in health and aging in CR mice fed diets that differ in fat composition. As occurs in other tissues and organs of mammals, kidney undergoes physiological and morphological changes during aging that lead to its deterioration and a consequent decline in renal function (Martin & Sheaff, 2007; Bolignano et al., 2014). Although CR had an important effect of decreasing serum urea content, no differences were found due to dietary fat. Also, creatinine clearance was similar in all experimental groups. These results are in accordance with those reported in rats as no striking changes in both metabolites were detected in aged ad libitum or CR-fed rats (Cui et al., 2012, 2013; Ning et al., 2013). On the other hand, we show here that p16 expression increased in CRS group during aging but maintained lesser values than those obtained for CON animals, which is in accordance with the results reported by other authors (Cui et al., 2012, 2013). Diet lipid composition did not impact age-related changes in p16 expression in CR mice.

Morphological and biochemical changes in glomeruli and PCT have been described in aged kidney. These changes include increased glomerulosclerosis, podocyte loss, GBM thickening, accumulation of abnormal mitochondria, and alteration in autophagy (Lindeman & Goldman, 1986; McKiernan et al., 2007; Bolignano et al., 2014). Most
of these changes have been observed in our samples in all of the dietary groups, but especially in the CON animals. However, differences were found in CR groups depending on the dietary fat.

Although the proportion of sclerotic glomeruli was similar in CON and CRL mice after 18 months of intervention, we found pronounced changes in nonaltered glomeruli which affected the glomerular basement membrane (GBM) thickness, the separation between contiguous podocyte processes (filtration slits, FS) and podocyte foot processes (PFP) width in the zone of contact with the GBM. These structures are essential components of the renal filtration barrier. Strikingly, GBM thickness increased after 6 months of CR when soy or fish oil were the primary dietary fats in comparison with CRL animals. However, 18 months of CR resulted in a remarkable increase of this parameter in all of the experimental groups. Wiggins et al. (2005) found similar results in rats subjected to CR and suggested that the increase in GBM thickness is an age-associated phenomenon and largely unrelated to diet. Although our results partially fit with these observations, it was noticeable that GBM thickness did not increase to the same extent in all of CR groups, with CRL-fed mice showing lesser values compared to all other CR groups, indicating that under CR dietary fat may partially prevent the increase in GBM thickness during aging.

Filtration slits width decreased during aging in control mice, a phenomenon that was prevented in CRS-fed animals. Moreover, our results show a negative correlation between GBM thickness and FS width in our animal model. On the other hand, it has been shown that aging induces expansion of podocyte processes (Wiggins et al., 2005; Hartleben et al., 2010), a phenomenon that could result in the reduction of FS width. In our control animals, aging also induced podocyte processes expansion, a phenomenon that was partially prevented by CR. However, dietary fat greatly affected this parameter. Thus, PFP width increased in aged CRF mice but remained unaltered in CRL-fed animals. Furthermore, our results show a negative correlation between PFP and FS width indicating that the narrowed FS found during aging can be due to PFP expansion. We also found a positive correlation between PFP width

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Aging also affected mitochondrial morphology and mass in PCT cells from mice submitted to CR. With the exception of CRL-fed mice, aging resulted in increased mitochondrial volume and decreased circularity. These results were especially prominent in 18-month CRS and CRF mice in which mitochondrial volumes increased nearly 45% in comparison with their younger counterparts. These results are in line with those obtained for mitochondrial Nv and the expression levels of PGC-1α, the key master of mitochondrial biogenesis regulation, and its downstream targets NRF1 and TFAM. Thus, low expression of these proteins was found in those groups showing enlarged mitochondria and low Nv values (CRS and CRF), and higher expression levels were detected in CRL group in which we found the smallest mitochondria and the highest Nv value. These results together with those concerning cellular and nuclear size, likely indicate differential adaptation mechanisms to the conditions imposed by CR. Interestingly, the response of PCT cells (in terms of cell and nuclear size and mitochondrial mass) to the different dietary fats under CR described here was similar to that found in mice hepatocytes (Khrawesh et al., 2013, 2014).

In an attempt to link glomerular ultrastructural changes with changes in epithelial PCT cells, several analyses were performed. Thus, we found a negative correlation between GBM thickness and mitochondrial mass (Vv and Nv) in PCT cells in such a way that a thick GBM corresponded to less mitochondrial mass in PCT cells. A similar negative correlation was also found when comparing FS width and Vv in PCT cells. These results seem to point out an adaptive response of PCT cells to changes in renal glomerular structures imposed by aging.

One of the hallmarks of aging is the generation of damaged organelles and molecular aggregates which may be removed by autophagy, and the decline in autophagic capacity is involved in the development of age-related diseases (see, for example, Rajawat et al., 2009). In kidney, decreased autophagy has been related to glomerulosclerosis, tubular atrophy, and interstitial fibrosis, and two different renal cell types have been reported to display autophagic activity depending on the physiological conditions: podocytes and PCT cells (Hartleben et al., 2010; Kume et al., 2010). In rodent kidney, aging has been shown to decrease phagocytic activity, a phenomenon that can be partially reverted by CR (Kume et al., 2010; Cui et al., 2012, 2013). In our model, Beclin-1, a protein involved in the control of autophagosome formation, increased during aging in CON animals. Among CR mice, Beclin-1 remained unaltered with age except in the CRF group in which a significant decrease was found.

The ratio of LC3-II/LC3-I has been shown to be an effective marker of autophagic flux. In our study, aging had no effect on this parameter in CON animals but markedly increased after 6 months of CR and these results seem to point out an adaptive response of PCT cells to changes in renal glomerular structures imposed by aging. However, further studies will be necessary to elucidate the physiological significance of this relationship. Also, the effects of specific dietary fats on GBM, FS, and PFP sizes have not been previously reported, and the results of the present study point out a possible role of dietary fat in the maintenance of these structures under CR conditions.

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The ratio of LC3-II/LC3-I has been shown to be an effective marker of autophagic flux. In our study, aging had no effect on this parameter in CON animals but markedly increased after 6 months of CR and these results are in accordance to those reported by several authors (Kume et al., 2010; Cui et al., 2012; Ning et al., 2013). Long-term CR did not induce additional increase of LC3 ratio in CRS group, although this value remained higher than that found in old CON mice. When comparing the different dietary fats in CR mice, we found the highest LC3 ratios in CRS group for both 6 and 18 months of CR. These results seem to indicate differential regulatory mechanisms of autophagy during calorie restriction depending on the fat source.

At the electron microscopy level, we found typical figures of autophagy in podocytes and PCT cells, especially in old animals.
However, enlarged structures consisting of multiple concentrically arranged electron-dense lamella occupying a high proportion of cellular volume were mainly found in PCT cells from old control and CR-fed mice. Nearly identical structures have been reported in a C57BL/6 mouse model of accelerated renal senescence (Yumura et al., 2006) and in other murine models with defective lysosomal activity (Porubsky et al., 2014). These authors identify these structures as lysosomes with accumulated lipofuscin and perhaps other nondegradable pigments, an idea compatible with the well-known fact that PCT cells may accumulate lipofuscin during aging (Melk et al., 2003). Furthermore, it has been proposed that a positive correlation exists between damaged mitochondria, lipofuscin accumulation and aging (Brunk & Terman, 2002). Due to its role in solute reabsorption, PCT cells show a high number of mitochondria making feasible the presence of these structures in this cell type as a consequence of altered autophagy or mitophagy processes for mitochondrial renewal. As PCT cells show a high rate of turnover (Fougeray & Pallet, 2015), it is not possible to assess whether proximal tubules showing an elevated number of altered lysosomes are fated to loss or regeneration.

As dietary lipids (lard, soybean oil, and fish oil) used in this and in previous studies are complex, comparisons between dietary lipids can be hampered by the fact that the lipids differ in multiple fatty acids. Thus, it cannot be unequivocally concluded which specific fatty acids were responsible for the biochemical and structural differences observed between CR groups. Nevertheless, we have previously shown that mitochondrial phospholipid fatty acid composition was altered in liver and skeletal muscle from CR mice in a manner that reflected the unsaturated fatty acid composition of the diet with the consequent increase of n-3 and n-6 fatty acids in CRF- and CRS-fed animals, respectively, and probably changing several properties of the membranes (Chen et al., 2012, 2013). On the other hand, CRL-fed animals showed a significantly higher proportion of mitochondrial monounsaturated fatty acids (especially oleic acid), a result that was accompanied by improved mitochondrial functions and ultrastructure (see Villalba et al., 2015 for a recent review). Thus, it is very likely that an increase in monounsaturated fatty acids such as oleic acid may be involved in the beneficial effect of lard as a dietary fat in CR-fed animals. However, further studies with purified fatty acids will be required to identify the specific fatty acids which influence health and lifespan in CR mice.

In summary, in this paper we report that long-term CR partially prevents or delays the appearance of several structural hallmarks of aging kidney, such as enlargement of GBM and PFP, FS narrowing, or PCT cells modification. However, these effects differed depending on the dietary fat. CRL mice showed an improved preservation of several renal structures (GBM thickness, PFP width, mitochondrial mass, size and shape, and autophagic processes in PCT cells) compared to other diet groups. These results fit well with those reported by López-Domínguez et al. (2015a) in which CR using lard as fat source resulted in extended longevity in comparison with other dietary fat (soy and fish oil), reinforcing the idea that dietary fat may have a crucial role in the determination of CR-mediated healthy aging in mice.

**Experimental procedures**

**Animals and diets**

A cohort of 64 ten-week-old male C57BL/6 mice was used (Charles River Laboratories, Wilmington, MA, USA). Mice were bred and raised in a vivarium at the Centro Andaluz de Biología del Desarrollo (CABD, Seville, Spain) under a 12-h light/dark cycle (8:00 a.m.–8:00 p.m.) and at controlled conditions of temperature (22 ± 3 °C) and humidity. The mice were fed a commercial rodent chow diet (Harlan Teklad #7012, Madison, WI, USA) for 14 days, and then, the animals were randomly assigned into four dietary groups and were fed a modified AIN-93G purified diet. The control group was fed 95% of a predetermined ad libitum intake (12.5 kcal). This slight restriction in food intake was initiated to prevent excessive weight gain during the study. The three CR dietary groups were maintained on 60% of the daily allowance of the control intake (8.6 kcal), and these diets were identical except for dietary lipid sources. The diets (percent total kilocalories per day) contained 20.3% protein, 63.9% carbohydrate, and 15.8% fat. The dietary fat for the control group was soybean oil. Dietary fats for the three CR groups were soybean oil (high in n-6 PUFAs, Super Store Industries, Lathrop, CA, USA), fish oil (high in n-3 PUFAs: 18% EPA, 12% DHA, Jedwards International, Inc. Quincy, MA, USA), or lard (high in saturated and monounsaturated fatty acids, ConAgra Foods, Omaha, NE, USA). To insure adequate linoleic acid levels, the CR-fish group was supplemented with soybean oil. Fatty acid composition of the dietary lipids has been detailed in a separate publication (Chen et al., 2012). All mice were housed individually and were fed with control or CR diets for 6 or 18 months, respectively. Filtered and acidified water was available ad libitum for all groups, and food was replaced every day between 8:00 and 9:00 a.m.

At either 6 or 18 months of CR, the animals were weighed and sacrificed by cervical dislocation after fasting O/N (or 12 h). Kidneys were quickly dissected and processed for ultrastructural analysis, and homogenates were also prepared for protein expression studies. Blood was collected by cardiac puncture just after cervical dislocation. Serum was obtained by centrifugation in Vacuette Z serum Sep Clot activator tubes for 10 min at 3000 g and stored in small aliquots kept at −80 °C until the determination of different blood metabolites. Serum creatinine was determined using an ELISA KIT (Alpha diagnostic international, San Antonio, Texas, USA) as indicated by the manufacturer. Urea was determined using the Reflotron plus system (Roche, Basel, Switzerland). All experimental procedures and animal handling were in accordance with the Pablo de Olavide University Ethical Committee rules, and the 86/609/EC Directive on the protection of animals used for experimental and other scientific purposes.

**Electron microscopy, planimetric and stereological analysis**

Small pieces from renal cortex were fixed and embedded in epoxy resin by conventional methods (see Data S1). The blocks were sectioned to obtain semi-thick (0.5–1 μm thickness) and thin (40–60 nm) sections. In semi-thick sections, we analyzed renal glomeruli morphology and thin sections were viewed and photographed for other measurements. GBM thickness, filtration slits (FS), and podocyte foot processes (PFP) width were measured using the ImageJ software (N.I.H.; Bethesda, MD, USA). Planimetric mitochondrial measurements of PCT cells were performed on pictures containing whole cells (see Figs S1 and S2) and using ImageJ software. From the same pictures, we obtained the mitochondrial stereological parameters Volume density (Vv) and Numerical density (Nv) using the semi-automatic application ‘WimStereology’ (Wimasis SL, Córdoba, Spain), based on a simple square lattice test system (Weibel, 1979). The relative number of autophagosomes and autophagic-related figures per cell surface area was also scored for each dietary group. Detailed information on ultrastructural procedures and applications is included in the Data S1 (Supporting information).

**Tissue processing for Western blotting analysis**

Kidneys were homogenized following a common protocol (see Data S1) to obtain properly samples for Western blot analysis. The samples were...
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Statistical analysis

Values were expressed as mean ± SEM. D’Agostino–Pearson tests were performed to determine data normality. The effect of CR was assessed by Student’s t-test (CRS group vs Control). In the case data did not pass the normality test, the nonparametric Mann–Whitney test was followed. The effect of dietary fat under CR was assessed by one-way ANOVA followed by a post hoc analysis (Tukey’s test for multiple comparisons) to assess significant differences among groups. Post hoc analysis of linear trend was also performed to investigate putative alterations of tested parameters among CR diets ordered as CRL→CRS→CRF, which resulted in a progressive increase of the n-6/n-3 ratio in phospholipid highly unsaturated fatty acids (Chen et al., 2012). In the case data did not pass the normality test, the nonparametric Kruskal–Wallis test was followed. Correlation analyses were performed by the nonparametric Spearman test.

Conflict of interest

None declared.

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Supporting Information

Additional Supporting Information may be found in the online version of this article at the publisher’s web-site.

Data S1 Supplementary methods.

Fig. S1 Cross-section of a proximal convoluted tubule (PCT) from a six-month old control animal.

Fig. S2 Representative images of cytoplasm portions of PCT epithelial cells from control (A) and 8-months CR-submitted animals with different dietary fats (B, CRL; C, CRS and D, CRF) showing a relatively large number of apoptosis (panels A, B and C) and glomerular structures in PCT cells (panel E). GBM thickness in PCT cells. In panel A, famotidine (F) versus control (C). In panels B–E, in podocytes foot processes (PFP) versus control (C). In panels A, B, C, D and E, in podocytes foot processes (PFP) versus contol (C).

Fig. S3 Representative gels stained with Ponceau S used to normalize quantifications of the different antibody bands shown in this paper.

Fig. S4 Correlation analyses between different glomerular filtration structures (panels A, B and C) and glomerular structures versus mitochondrial mass in epithelial cells from proximal convoluted tubules (D, E and F). Panel A shows filtration slits (FS) versus glomerular basal membrane (GBM) thickness; panel B, podocyte foot processes (PFP) versus GBM and panel C, PFP versus FS. Panel D depicts GBM thickness versus mitochondrial volume density (VV) in PCT cells; panel E, GBM thickness versus mitochondrial numerical density in PCT cells and panel F, PFP width versus mitochondrial VV in PCT cells. In panel A, P < 0.001; in panels B–E, P < 0.05. In this figure C is CON and L, S and F are CRL, CRS and CRF respectively. Number 6 and 18 indicates the duration of dietary intervention period.