Folate Acts in *E. coli* to Accelerate *C. elegans* Aging Independently of Bacterial Biosynthesis

**Graphical Abstract**

**Highlights**

- Limiting folate in *E. coli*, not in *C. elegans*, increases worm lifespan
- An *E. coli* screen for worm longevity identifies folate synthesis as a target
- Folate synthesis influences *E. coli* physiology independently of growth
- Bacterial folate synthesis may be a sustainable target for chronic disease

**Authors**

Bhupinder Virk, Jie Jia, Claire A. Maynard, ..., Noel Helliwell, Marta Cipinska, David Weinkove

**Correspondence**

david.weinkove@durham.ac.uk

**In Brief**

Virk et al. show that inhibiting *E. coli* folate synthesis does not increase *C. elegans* lifespan through changes to *C. elegans* folate, but by acting through an *E. coli* activity. They find nine *E. coli* mutants that extend lifespan. The gene identities suggest that mild bacterial toxicities accelerate host aging.
Folate Acts in *E. coli* to Accelerate *C. elegans* Aging Independently of Bacterial Biosynthesis

Bhupinder Virk,1,7,8 Jie Jia,1,2,4,7 Claire A. Maynard,1,7 Adelaide Raimundo,1 Jolien Lefebvre,1,5 Shane A. Richards,1 Natalia Chetina,1 Yen Liang,1 Noel Helliwell,1 Marta Cipinska,1,6 and David Weinkove1,6,*

1School of Biological and Biomedical Sciences, Durham University, South Road, Durham DH1 3LE, UK
2Department of Nutrition, Shanghai Jiao Tong University School of Medicine, Shanghai 200025, China
3Key Laboratory of Pediatric Gastroenterology and Nutrition, Shanghai Institute for Pediatric Research, Shanghai 200092, China
4Department of Clinical Nutrition, Xin Hua Hospital affiliated to SJTU School of Medicine, Shanghai 200092, China
5Department HIVB, VIVES, Wilgenstraat 32, 8800 Roeselare, Belgium
6Biophysical Sciences Institute, Durham University, South Road, Durham DH1 3LE, UK
7Co-first author
8Present address: Bioinformatics Department, The Babraham Institute, Cambridge CB22 3AT, UK
*Correspondence: david.weinkove@durham.ac.uk
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SUMMARY

Folates are cofactors for biosynthetic enzymes in all eukaryotic and prokaryotic cells. Animals cannot synthesize folate and must acquire it from their diet or microbiota. Previously, we showed that inhibiting *E. coli* folate synthesis increases *C. elegans* lifespan. Here, we show that restriction or supplementation of *C. elegans* folate does not influence lifespan. Thus, folate is required in *E. coli* to shorten worm lifespan. Bacterial proliferation in the intestine has been proposed as a mechanism for the life-shortening influence of *E. coli*. However, we found no correlation between *C. elegans* survival and bacterial growth in a screen of 1,000+ *E. coli* deletion mutants. Nine mutants increased worm lifespan robustly, suggesting specific gene regulation is required for the life-shortening activity of *E. coli*. Disrupting the biosynthetic folate cycle did not increase lifespan. Thus, folate acts through a growth-independent route in *E. coli* to accelerate animal aging.

INTRODUCTION

Both nutrition and the host-associated microbiota are thought to impact longevity (Heintz and Mair, 2014; Rizza et al., 2014). Diet influences the metabolism of gut microbes, which in turn can synthesize nutrients for the host. These interactions make it difficult to unravel the contributions of diet and the gut microbiota to long-term health (Lozupone et al., 2012). This complexity can be addressed with model systems such as the nematode *Caenorhabditis elegans* (Collins et al., 2008). Yet even here, there are numerous interactions between the nutrient agar medium, the *Escherichia coli* bacterial lawn, and the worm. Chemical manipulations of the medium and genetic manipulations of both *E. coli* and *C. elegans* provide tools to understand these interactions (Weinkove, 2015).

Folates in their reduced tetrahydrofolate (THF) form are required as enzymatic cofactors in the folate cycle; a series of metabolic steps found in all cells (including both bacteria and animals) required for cell biosynthesis. Products include purines, pyrimidines, glycine, and methionine, which are required to generate the methyl donor molecule S-adenosyl methionine (SAM) (Bailey and Gregory, 1999). Animals cannot synthesize folates and so obtain folates from their diets and associated microbes (Asrar and O’Connor, 2005; Lakoff et al., 2014). Our previous research showed that *C. elegans* lifespan is increased when *E. coli* folate synthesis is disrupted either by a mutation in the gene aroD, which is needed to make aromatic compounds including the folate precursor para-aminobenzoic acid (PABA), or by sulfamethoxazole (SMX), a sulphonamide drug that competes with PABA for the active site of the enzyme dihydropteroyl synthase (Virk et al., 2012). This enzyme is a key step in folate biosynthesis and is absent from animals. *C. elegans* obtains folates from *E. coli* and thus several possible mechanisms might explain why *E. coli* folate synthesis affects *C. elegans* lifespan. Distinguishing the effects of folates in bacteria and folates in their animal hosts is important because folate supplementation is beneficial to human health and any intervention would need to maintain healthy levels of serum folate.

Dietary, or caloric, restriction has been shown to extend the lifespan of *C. elegans* (Greer and Brunet, 2009; Mair and Dillin, 2008). SMX does not slow *E. coli* growth and therefore has no effect on food availability. Furthermore, *C. elegans* grow and reproduce normally (Virk et al., 2012). Thus, a limitation of macronutrients is an unlikely explanation. Alternatively, inhibition of *E. coli* folate synthesis may influence *C. elegans* lifespan by limiting dietary folate and/or a specific change in folate-dependent nutrients (Lee et al., 2015). For example, restriction of methionine increases lifespan in rodents and influences lifespan in *Drosophila* (Grandison et al., 2009; Sanchez-Roman and Barja, 2013). Mutation of *C. elegans* sams-1, the gene encoding SAM synthase, extends lifespan (Hansen et al., 2005).
The diabetes drug metformin increases *C. elegans* lifespan in a manner dependent on the *E. coli* strain and changes in *C. elegans* folate and methionine metabolism are implicated in mediating the lifespan extension (Cabreiro et al., 2013).

Another possible explanation is that folate synthesis inhibition increases *C. elegans* lifespan by altering *E. coli* physiology. *E. coli* can accumulate in the intestine of older *C. elegans* adults and because treatment of *E. coli* with antibiotics or UV increases worm lifespan, this accumulation is widely thought to accelerate *C. elegans* aging (Garigan et al., 2002; Gems and Riddle, 2000; McGee et al., 2011). More subtly, changes in bacterial toxicity caused by changes in bacterial metabolism might influence *C. elegans* because treatment of *C. elegans* with antibiotics or UV increases worm lifespan by influencing bacterial respiration rather than dietary intake of Q (Saiki et al., 2008).

Here, we show that modulating folate uptake or the folate cycle in *C. elegans* does not affect lifespan, suggesting *E. coli* folate influences *C. elegans* lifespan by acting on *E. coli* physiology. Apart from the Q synthesis genes and *aroD* (Saiki et al., 2008; Virk et al., 2012), little is known about how *E. coli* genetics influences *C. elegans* lifespan. A genetic screen of over 1,000 *E. coli* mutants shows that bacterial growth does not correlate with *C. elegans* survival and only a few specific interventions increase *C. elegans* lifespan, including the mutation of genes involved in *E. coli* folate synthesis, but not in the *E. coli* folate cycle. In addition to its role in bacterial growth, we propose that folate acts to change *E. coli* physiology in a way that accelerates *C. elegans* aging.

**RESULTS**

**Genetic Disruption of *C. elegans* Folate Uptake and Restoration by Supplementation**

To distinguish the effects of *C. elegans* folate from *E. coli* folate, we targeted folate uptake in *C. elegans*. The reduced folate carrier FOLT-1 takes up THFs across the intestinal epithelial membrane (Balamurugan et al., 2007). The published *folt-1* deletion allele causes sterility, so we turned to another *C. elegans* mutant predicted to disrupt folate uptake. In mammals, glutamate carboxypeptidase II (GCPII) cleaves glutamates from polyglutamated THFs in the gut, to create monoglutamated folates that are preferentially imported by folate carriers and transporters (Halsted et al., 1998). The *E. coli* diet contains predominantly polyglutamated THFs (Kwon et al., 2008; Virk et al., 2012), suggesting that *C. elegans* requires GCPII activity. There are three *C. elegans* genes that encode a GCPII homolog (Supplemental Information). Animals lacking the GCPII gene *gcp-2.1, WB Gene: WBGene00020082*, appear healthy and develop normally on *E. coli* OP50, but develop slowly and are uncoordinated and sterile on OP50 treated with 128 μM SMX (Figure 1A). This phenotype is rescued completely by supplementation with 10 μM formyl THF monoglutamate, a naturally occurring reduced folate also known as folic acid or leucovorin (Figures 1A and 1B). Folic acid can rescue the *gcp-2.1* phenotype at a 20-fold lower concentration than can folic acid, the oxidized folate used commonly in dietary supplements (Figure 1B). These results are consistent with the specificity of FOLT-1 for reduced folates (Balamurugan et al., 2007) and a role for GCP-2.1 in folate uptake (Figure 1C).

*E. coli* does not have folate uptake transporters, but can synthesize folate from the folate breakdown products PABA, which can freely diffuse through membranes, and PAB-Glutamate, which is taken up by an active transport system (Hussein et al., 1998). Thus, folic acid restores folate synthesis in an *aroD* mutant, most clearly at concentrations of 100 μM or more (Virk et al., 2012). In the presence of SMX, which competes with PABA for the enzyme dihydrofolate synthase, it is likely that more folate breakdown products would be required to restore folate synthesis. For example, PABA is required at a concentration of 250 μM to fully reverse the lifespan extension caused by 505 μM (128 μg/ml) SMX (Virk et al., 2012). It is unlikely that 10 μM folic acid would provide enough breakdown products to restore *E. coli* folate synthesis in competition with 505 μM SMX. Thus, we conclude that folic acid rescues *gcp-2.1* by directly supplementing *C. elegans* folate (Figure 1C).

**Modulation of *C. elegans* Folate Status Does Not Influence Lifespan**

Turning to effects on aging, we found that the *C. elegans gcp-2.1* mutant has a similar, if not slightly shorter lifespan, than wild-type controls (p = 0.0227; Figure 2A). Furthermore, the increased *C. elegans* lifespan caused by SMX was unaffected by supplementation with 10 μM folic acid (Figure 2B). Methotrexate (MTX), a dihydrofolate reductase inhibitor specific to animals, inhibits the *C. elegans* folate cycle. 100 μg/ml MTX causes developmental defects in *nuc-1* mutants and is five times the concentration required to cause this phenotype (Mello et al., 1991; Virk et al., 2012). MTX did not affect the lifespan of *C. elegans* and also failed to influence the lifespan of worms on SMX-treated bacteria (Figure 2C). Together, these results suggest that the SMX-induced lifespan increase cannot be explained by decreased *C. elegans* folate uptake or impaired folate-dependent *C. elegans* metabolism.

**SMX and Kanamycin Treatment Cause an Identical Increase in *C. elegans* Lifespan**

Treating *E. coli* with ultraviolet irradiation or antibiotics increases *C. elegans* lifespan, suggesting that *E. coli* possess a life-shortening activity (Garigan et al., 2002; Gems and Riddle, 2000). We compared OP50 treated with SMX, which does not influence *E. coli* viability, with OP50 treated with kanamycin, a bacterial translation inhibitor. Addition of kanamycin to the bacterial lawn stops cells forming further colonies (Virk et al., 2012). Using large cohorts, worms maintained on kanamycin-treated bacteria showed an almost identical survival curve to those on bacteria treated with SMX (25% increase in mean lifespan compared to wild-type, p = 0.0001), with a small further increase in lifespan (4%, p = 0.0008) when both drugs were combined (Figure 2D). While there are many possible explanations for this result and SMX and kanamycin have very different targets and effects on *E. coli* metabolism, the lack of a substantial additive effect suggests that both drugs might inhibit a shared downstream process that shortens *C. elegans* lifespan.
It Is Unlikely that *E. coli* Shortens Lifespan Solely through Intestinal Accumulation

Kanamycin and similar treatments are thought to increase *C. elegans* lifespan by preventing *E. coli* accumulation in the intestinal lumen (Garigan et al., 2002). To test whether SMX prevents accumulation, we performed lifespan experiments with worms maintained on untreated OP50 *E. coli*, but SMX treatment delays the growth of *gcp-2.1* mutants, and this defect can be rescued by 10 μM folic acid. The images were taken after 48 hr of growth at 25°C. (B) Quantification of the growth, as measured by body length. The error bars represent SD. The *gcp-2.1* mutant growth on SMX is restored at 5 μM and 10 μM folic acid and partially with 1 μM folic acid (*p < 0.05, **p < 0.01, and ***p < 0.005*) (t test). Folic acid can fully rescue growth only at 250 μM.

(C) Model showing that *E. coli* folate synthesis generates THFs with up to eight glutamate residues and various one carbon groups, xTHFGlu_{1–8} (x = methyl, formyl, methenyl, and methylene). The *C. elegans* GCPII GCP-2.1 cleaves glutamate residues from xTHFGlu_{2–8} to generate xTHFGlu_{1}, the preferred substrates of the *C. elegans* reduced folate transporter FOLT-1. Folinic acid, as an xTHFGlu_{1} (5-formyl THF), can be taken up directly by FOLT-1, bypassing GCP-2.1.
that these bacteria are capable of producing increased protein. SMX prolonged the time that C. elegans stay mobile, consistent with a decrease in the rate of aging (Figure 3Q). The structure and function of the C. elegans intestine declines with age. Thus, like motility, bacteria accumulation is a biomarker, but not necessarily a cause, of aging.

**E. coli Mutants Influence C. elegans Lifespan Independently of E. coli Growth**

To further understand how *E. coli* influence *C. elegans* lifespan, we conducted a screen of over 1,000 *E. coli* K12 mutants from the Keio collection (Baba et al., 2006). We tested deletions in all non-essential genes predicted to encode enzymes in the folate cycle or related pathways (25 genes; Table S2) and deletions of 981 randomly selected genes of known function (Experimental Procedures; Table S2). Scoring across the lifespan is impractical for a large-scale screen, so we scored survival at a single time point (day 11/12) close to the median lifespan. This strategy allows greater statistical power than scoring near the end of the survival curve and the identification of strains that shorten, as well as extend, *C. elegans* lifespan.

The mutants were scored in batches. Each batch contained several mutants and three wild-type strains. The distribution in survival shown by these controls did not differ from the distribution across all mutants tested, suggesting that mutating single *E. coli* genes had no detectable large-scale effect on *C. elegans* lifespan (Supplemental Information; Figure S2). To account for batch-to-batch variation, we subtracted the mean survival of strains that increased *C. elegans* lifespan (Figure S2). Using growth data from Baba et al. (2006), we found no correlation between strain growth and survival of *C. elegans* (Figure 4A), suggesting that *E. coli* growth rate does not influence *C. elegans* aging.

**Nine *E. coli* Mutants Robustly Increase *C. elegans* Lifespan**

To be confident of identifying individual mutants that increased *C. elegans* lifespan, we repeated the screen for the 67 strains that caused worms to survive at least 15% more than on the control strain. We undertook full lifespan analysis of the 22 strains that passed this second round. Finally, we retested the 11 strains that passed this third round and included full lifespan analysis of the strains with the mutation complemented by the wild-type *E. coli* gene. This step ruled out lifespan increases from spontaneous second-site mutations (Supplemental Information). There were nine deletions that passed this final test, representing less than 1% of genes tested.

Three of the identified mutants (*metL*, *ihfA*, and *ihfB*) caused the bacterial lawn to appear more liquid than normal. This environment would have a strong influence on *C. elegans* physiology (Lewis and Fleming, 1995). The other mutants discovered had no visible effect on the *E. coli* lawn and so are more likely to influence lifespan through biological, rather than physio-chemical mechanisms. These mutants included a deletion of *pcoS*, a stationary phase sigma factor. This transcription factor regulates over 200 genes in response to low nutrients or other stresses (Battesti et al., 2011). Other deletions that extended lifespan included *tatC*, a gene encoding a component of the twin arginine translocation pathway, which transports folded proteins to the periplasm (Stanley et al., 2001), *ompA*, an abundant constituent of the outer membrane (Smith et al., 2007), and *znuB*, which encodes part of the ZnuABC zinc transporter, needed to take up zinc at low concentrations (Patzer and Hantke, 1998). Finally, we isolated deletions in *pabA* and *pabB* (Figure 4B). These genes encode two enzymes that associate and catalyze key steps in PABA synthesis (Green et al., 1996). In summary, the screen underlined the importance of *E. coli* folate synthesis in *C. elegans* lifespan regulation and identified other genes with diverse functions.

Apart from the *pabA* and *pabB* mutants, the mutants grow slower on the petri dish than the wild-type strain (Figure S3A). There was a positive, but not significant, correlation between extent of the lifespan extension and the growth rate of the strain.
We noticed that *C. elegans* spent more time on the bacterial lawn if the *E. coli* strain was one of the life-extending mutants or had been treated with SMX (Figure S3C). SMX treatment of OP50 also produced a similar decrease in aversion. Thus, the increased lifespan of *C. elegans* is not caused by decreased exposure to food or to *E. coli*. *C. elegans* avoids toxic bacteria, and this avoidance is thought to be triggered by perception of disruption to worm metabolism caused by bacterial toxins (Melo and Ruvkun, 2012). Thus, the mutations isolated in the screen, or chemical inhibition of folate synthesis, may remove toxicity from *E. coli*.

Disruption of the *E. coli* Folate Cycle Does Not Increase *C. elegans* Lifespan

Although the identification of *pabA* and *pabB* was consistent with *E. coli* folate synthesis inhibition increasing *C. elegans* lifespan, we were surprised that no genes involved in the folate cycle or related pathways were identified. An exception was *metL*, which encodes an enzyme needed in two early stages of methionine biosynthesis. However, deletions in other methionine synthesis genes did not have a lifespan phenotype. Because of the variation found in the screen, we wanted to make sure that we had not missed any folate-related genes.

**Figure 3. Intestinal Accumulation of Bacteria Does Not Occur in All Animals and Is Not Prevented by SMX**

(A) Accumulation in recently dead animals as assessed by visualizing *E. coli* GFP in the intestinal lumen. The data are pooled from two biological replicates.

(B) Numbers of alive worms at indicated time points with classification of accumulation.

(C) Motility analysis of *glp-4(bn2)* worms on OP50 treated with 0, 16 μg/ml, and 128 μg/ml SMX. Each worm was scored as belonging to motility class A (constantly moving), B (moves when prodded), or C (twitches only) as described (Herndon et al., 2002).
We performed full lifespan analysis on *E. coli* mutants in 23 non-essential genes involved in the folate cycle or related metabolic pathways. None of these mutants extended lifespan apart from a small, but significant, effect of a deletion of *glyA* (Table S1). Thus *E. coli* folate synthesis, but not the folate cycle, which is needed for *E. coli* growth, limits *C. elegans* lifespan.

We tested how modulating *E. coli* folate synthesis influenced *C. elegans* lifespan by maintaining worms on the *pabA* mutant several generations of subculture in the defined medium. The *pabA* mutant was unable to grow without the addition of PABA. Growth was restored by as little as 50 nM PABA, whereas 100 μM folic acid only incompletely rescued growth (Figure S4A).

*C. elegans* Lifespan Can Be Altered through the Availability of PABA to *E. coli*

Folate synthesis is essential for *E. coli* growth because *E. coli* cannot take up intact folate. However, SMX or mutation of *pabA* or *pabB* do not decrease *E. coli* growth rate under the conditions that they increase *C. elegans* lifespan (Virk et al., 2012) (Figure S3A). Thus, these interventions must only remove folate in excess of that required for *E. coli* growth. To understand the relative levels at which folate synthesis is required to limit lifespan compared to requirements for growth, we took advantage of the fact that PABA synthesis in *E. coli* can be bypassed by adding exogenous PABA, which can diffuse across membranes and that *C. elegans* cannot make folate from PABA. The growth of *pabA* and *pabB* mutants on peptone-based NGM, suggests this medium contains a source of PABA. To remove this PABA while minimizing changes to the nutritional conditions, we replaced peptone with a defined mix of amino acids based on the composition of peptone, an undefined digest of soy protein (Supplemental Information). When subcultured from the rich lysogeny broth (LB) broth, the *pabA* and *pabB* mutants grow well on this defined medium. However, after several generations of subculture in the defined medium, the *pabA* mutant was unable to grow without the addition of PABA. Growth was restored by as little as 50 nM PABA, whereas 100 μM folic acid only incompletely rescued growth (Figure S4A).
grown on defined medium supplemented with a range of PABA concentrations. Before seeding, the pabA mutant was cultured in liquid defined medium containing 0.2 μM PABA. Subsequent growth of the pabA and wild-type bacterial lawn was the same at all PABA concentrations (Figure S4B). However, on medium supplemented with 0.1 μM PABA, C. elegans on the pabA mutant lived 39% longer than worms on wild-type bacteria (p = < 0.0001), or 50% longer than worms on pabA bacteria supplemented with 1 μM PABA (p = < 0.0001; Figure 4C). Supplementation of pabA E. coli with 0.2 μM PABA gave an intermediate result (28% increase, p = 0.0009, compared to 1 μM PABA). Increasing the concentration to 100 μM PABA did not decrease lifespan (Figure 4C) on either pabA or the wild-type bacteria, suggesting that PABA is not toxic. Rather, C. elegans lifespan is increased when excess folate synthesis is removed.

**DISCUSSION**

**C. elegans Folate and C. elegans Lifespan**

Inhibiting *E. coli* folate synthesis decreases *C. elegans* folates (Virk et al., 2012), but we have shown that these changes in *C. elegans* folate are not responsible for the increased lifespan. Lifespan is also unaffected by *C. elegans* folate cycle inhibition, by MTX (Figure 2C), or by supplementation of vitamin B12, which is required for a key step in the folate cycle (Watson et al., 2014).

**Intestinal Accumulation of *E. coli*: a Cause or Consequence of *C. elegans* Aging?**

Our analysis of accumulation of *E. coli* in the intestines of individual animals (Figure 3) does not exclude accumulation as the mechanism by which folate synthesis influences lifespan, but our data are also consistent with intestinal accumulation of *E. coli* being a consequence, rather than a cause, of intestinal aging. The intestine loses its structure early in aging worms (Hern don et al., 2002; McGee et al., 2011) and an age-related delay in the passage of *E. coli* caused by decreased pharyngeal pumping, decreased defecation, or breakdown in lumenal structure, would likely lead to bacterial accumulation. To our knowledge, it has never been demonstrated that preventing bacterial accumulation increases *C. elegans* lifespan. Electron microscopy studies have failed to find evidence of *E. coli* OP50 invasion in the *C. elegans* intestine (McGee et al., 2011) (David Hall, personal communication) and strains of *E. coli* used in *C. elegans* experiments lack the O-antigen needed for invasive pathogenesis (Browning et al., 2013). Through increased contact with intestinal cells, accumulation in the intestine may enhance other mechanisms by which *E. coli* are pathogenic, thus accelerating a cycle of functional loss in the aging intestine.

Second, we uncoupled the effects of *E. coli* growth from the *E. coli* activity that accelerates aging in *C. elegans*. Though SMX does not slow *E. coli* growth, we suggest it prevents a process that is also prevented by kanamycin. This process might be a factor that is induced by high folate levels and requires kanamycin-sensitive translation. The *E. coli* screen showed that lifespan is increased by only a few mutations, while many mutations slow *E. coli* growth without extending *C. elegans* lifespan. Thus, an alternative to the growth-dependent model is that a specific *E. coli* activity shortens lifespan, and this activity is blocked by treatment with kanamycin and other antibiotics.

**How *E. coli* Influences *C. elegans* Aging**

The comparison with kanamycin suggests that inhibiting folate synthesis removes a pro-aging activity rather than producing an anti-aging activity. A common factor of *E. coli* genes isolated in the screen such as *pabA, pabB, rpoS, tatC, znuB, and ompA*, is that they reduce virulence when mutated in a wide range of pathogenic bacteria (Brown and Stocker, 1987; Dong and Schellhorn, 2010; Gabbianelli et al., 2011; Ochsner et al., 2002; Teng et al., 2008). Lab strains of *E. coli* used for *C. elegans* culture do not have known virulence factors, but the genes isolated in the screen might regulate other, as yet unknown, factors with a milder, long-term effect on their hosts. The decreased aversion to *E. coli* mutants identified in the screen, or to *E. coli* treated with SMX, is consistent with the removal of a toxin (Figure S3C) (Melo and Ruvkun, 2012). It is likely that some of the many peptides and compounds secreted by *E. coli* are toxic to *C. elegans*. These products may or may not influence lifespan. This study suggests that *E. coli* shortens *C. elegans* lifespan through a form of toxin-based virulence that is milder than observed with human pathogens, but may nevertheless be important for chronic disease and aging.

It is interesting to note that many *E. coli* genes and processes were not found to increase *C. elegans* lifespan robustly in the screen. Surprisingly, genes involved in ubiquinone synthesis or respiration were not found (Saiki et al., 2008), although the Keio ubiG mutant did not grow sufficiently to be included in the screen. The *E. coli* genes of unknown function, which constitute over a third of the mutant collection (Saba et al., 2006), were not screened and may be important for interactions with the host by, for example, synthesizing as yet uncharacterized toxins.

**The Role of Bacterial Folate**

Our work suggests that in bacteria, folate has functions beyond its role in biosynthetic one-carbon metabolism. Mutation of *pabA* or *pabB* attenuates virulence in invasive bacteria, and the accepted explanation is that these mutants cannot grow in the absence of PABA (Brown and Stocker, 1987; Chimalapati et al., 2011). However, another explanation is that these mutants are less able to produce toxins. Likewise, sulfonamides, which are less effective than most antibiotics in stopping bacterial growth, may be effective by preventing toxicity rather than growth. RpoS is an *E. coli* sigma factor activated in stationary phase to coordinate a global stress response, which includes increased virulence. Of the hundreds of genes reported to be under RpoS transcriptional control, *pabA* in *E. coli* (Weber et al., 2005) and *pabB* in *Bacillus subtilis* (Eymann et al., 2002) have been implicated in microarray experiments. Thus, folate synthesis may be stimulated by RpoS activity.

**Possible Implications for Human Aging and Disease**

We have presented evidence that *E. coli* accelerates *C. elegans* aging independently of *E. coli* growth and *C. elegans* folate metabolism. If a similar relationship existed in the human gut, molecular characterization of this mechanism may uncover targets to
intervene in aging and chronic disease. Chronic conditions such as obesity and inflammatory bowel disease are characterized by a dysbiosis of the microbiota, leading to an overgrowth of gamma proteobacteria, such as *E. coli* (Winter and Bäumler, 2014). Interestingly, small intestine bacterial overgrowth leads to increased levels of serum folate, originating from dominating opportunistic bacteria (Camilo et al., 1996; Lakhani et al., 2008). Dysbiosis and the consequent overabundance of gamma proteobacteria may be an important factor in aging (Clark et al., 2015). This study shows that bacteria folate synthesis can be targeted without compromising host folate status, which can be maintained by selective supplementation.

**EXPERIMENTAL PROCEDURES**

**C. elegans Strains**

N2 (wild-type), SS104 *glp-4(bn2)*, UF208 (wild-type), and UF209 *gcp-2;1(ok1004). See Supplemental Information for *gcp-2;1* analysis.

**E. coli Strains**

OP50 *ura* (Brenner, 1974), OP50-GFP (containing plasmid pFPV25.1) (Labrousse et al., 2000), BW21153 (Keio collection wild-type), and WT Kan: BW21153 pGreen 0029 (Kan+). Strains from Keio collection as listed (Table S2). Keio strains transformed with complementation plasmids (Tables S1 and S3).

**Culture Conditions**

NGM was prepared using high purity agar as described (Virk et al., 2012). As necessary, 25 µg/ml carbenicillin was added to maintain plasmid selection (OP50-GFP and complemented Keio mutants). Kanamycin was added after 24 hr of bacterial growth as described (Garigan et al., 2002). Folinic acid, folic acid, and Mtx were from Schircks Laboratories. For the defined media, peptone was replaced by purified amino acids (Supplemental Information) and 0.2 x trace metals (Studier, 2009). Calcium chloride was omitted as it is in the trace metals.

**Lifespan Analysis**

Survival analyses were performed as described (Virk et al., 2012). Worms were cultured at 15°C and shifted to 25°C at the L3 stage. At 25°C, *E. coli* metabolism is more active than at lower temperatures (Virk et al., 2012). At the L4/young adult stage, animals were placed on bacteria under the experimental conditions. All lifespan data is in Table S1. Statistical significance was determined using log rank and Wilcoxon tests of the Kaplan-Meier survival model.

**GFP Bacterial Accumulation Assay**

Worms were prepared and set up as for lifespan analysis using OP50-GFP. From day 5, using a Leica M165 FL stereomicroscope with a GFP filter with a 510 nm + long pass emission spectrum, individual animals were scored every 2 days for survival and as having no accumulation, partial accumulation, or full accumulation (Figure S1).

**Motility Assay**

Animals were prepared as for lifespan analysis as described, with 16 plates per condition used and ten worms per plate. Animals were classified as described (Herndon et al., 2002).

**C. elegans Growth/Body Size Analysis**

Animals were imaged after 2 days of 25°C growth after a synchronized egg lay. Images were analyzed for body length as described in Supplemental Information.

**Screening Method**

See Supplemental Information.

**Gene Complementation of Keio Mutants**

The relevant genomic region was PCR amplified from BW21153 using primers with restriction sites (Table S3), cloned into the low copy pMMB67EH AmpR plasmid (Virk et al., 2012), and transformed into the relevant mutant strain. The plasmid was used as a control.

**Measuring *E. coli* Lawn Growth**

To quantify bacteria in bacterial lawns, 2 ml of M9 was added to a 6 cm plate. The lawn was scraped off with a glass scraper. The liquid was then removed to a fresh tube. The total volume of the removed liquid was multiplied by the OD600 after a 5-fold dilution to generate a relative measure of bacteria.

**Aversion Assay**

Aversion was measured as number of worms off the lawn/total number of worms as described in Melo and Ruvkun (2012).

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Supplemental Experimental Procedures, four figures, and three tables and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2016.01.051.

**AUTHOR CONTRIBUTIONS**


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**REFERENCES**


Supplemental Information

Folate Acts in *E. coli* to Accelerate *C. elegans* Aging Independently of Bacterial Biosynthesis

Bhupinder Virk, Jie Jia, Claire A. Maynard, Adelaide Raimundo, Jolien Lefebvre, Shane A. Richards, Natalia Chetina, Yen Liang, Noel Helliwell, Marta Cipinska, and David Weinkove
Supplemental Information
Figure S1. Distinguishing intestinal accumulation of *E. coli* OP50-GFP. Related to Figure 3. A) Representative images of single worms (observed using a long-pass green filter) illustrating the distinction between gut autofluorescence (yellow-green, yellow arrows) and GFP-expressing *E. coli* (bright green, green arrows). B) Representative images of single worms as classified according to extent of colonization by GFP-expressing *E. coli*. 
Figure S2. Estimating batch and plate variation on worm survival. Related to Figure 4. A) Histogram comparing the distribution in *C. elegans* survival in mutant and wild type strains. B) Normalizing the distribution to account for batch-to-batch variation: mutant survival ($S_m$) – wild type survival in the same batch ($S_w$). C) i) Variation in the number of WT worms scored per plate across batches ii) and the proportion of worms that survived in each plate. The dashed line indicates the maximum-likelihood fit for the overall survival fraction, $p$. All whiskers indicate the most extreme values. D) Best fitting beta distributions describing the variation in mean WT survival fraction to day 11 among plates (solid line), and variation in mean survival fraction between plates within a batch (dashed line). Here, we have depicted among plate variation for a batch having the overall mean survival, $p$. See Supplemental Experimental Procedures.
Supplemental Tables

(Excel Files)

**Table S1. Lifespan summaries. Related to Figures 2 and 4.** Contains conditions, numbers and relevant statistical analysis for all lifespan experiments in the study.

**Table S2 Keio collection strains included in the screen. Related to Figure 4.** A) Strains tested and their survival in the first round. B) Strains excluded because they did not grow on NGM plates.

**Table S3 Strains selected for 2nd, 3rd and 4th rounds of the screen and the results. Related to Figure 4.** Includes primers used to make complementation plasmids and the corresponding strains. Strains in red were added to 2nd and 3rd rounds because they were of interest.
Figure S3

A) Growth on NGM plates over 72 hours is significantly lower than wild type apart from *pabA* (no statistical difference), and *pabB* (significantly higher). See Experimental Procedures. Error bars represent standard deviation. B) Positive but not statistically significant correlation between bacterial growth on plates (A) and *C. elegans* lifespan increase for positive mutants from screen (Table S3). $R^2 = 0.06$, $P = 0.53$. C) Aversion of *C. elegans* strains to the bacterial lawn is decreased with the isolated mutants or with SMX. Error bars represent standard deviation. Aversion was statistically lower than control ($P < 0.01$) apart from on deletions of *pabB, ompA, tatC*, for which aversion is statistically lower only when data for N2 and SS104 are combined.

Figure S3. Analysis of the 9 mutants isolated in the screen. Related to Figure 4. A) Growth on NGM plates over 72 hours is significantly lower than wild type apart from *pabA* (no statistical difference), and *pabB* (significantly higher). See Experimental Procedures. Error bars represent standard deviation. B) Positive but not statistically significant correlation between bacterial growth on plates (A) and *C. elegans* lifespan increase for positive mutants from screen (Table S3). $R^2 = 0.06$, $P = 0.53$. C) Aversion of *C. elegans* strains to the bacterial lawn is decreased with the isolated mutants or with SMX. Error bars represent standard deviation. Aversion was statistically lower than control ($P < 0.01$) apart from on deletions of *pabB, ompA, tatC*, for which aversion is statistically lower only when data for N2 and SS104 are combined.
Figure S4. Growth of the *pabA* mutant. Related to Figure 4C and Figure 1. A) Growth curve of the *pabA* mutant in liquid defined media (Experimental Procedures) shows that 50 nM PABA is sufficient to restore bacterial growth. Folinic acid cannot completely restore growth at even at 100 μM. B) Growth on defined media plates with various concentrations of PABA after 72 hours, comparing *pabA* with WT under conditions used for Figure 4C. See Experimental Procedures. Error bars represent standard deviation. * p < 0.01 for growth on *pabA* being less than growth on WT Kan.
Supplemental Experimental Procedures

GCP-2.1 characterization and analysis. Related to Figure 1

Three predicted proteins in C. elegans are encoded by R57.1 (named gcp-2.1), C35C5.2 (named gcp-2.2) and C35C5.11 (named gcp-2.3). All show similarity to mammalian GCPII amino acid sequences. The gcp-2.1(ok1004) deletion mutant from the C. elegans knockout consortium was outcrossed 3 times using N2 to make UF209 gcp-2.1(ok1004). The wild type sibling strain UF208 was used as the control. To make the gcp-2.1 genomic transgene, a genomic fragment containing the predicted gcp-2.1 gene was amplified using the primers R57gen_5: CTTAGGTTGGATCTCGTTGCTTGC and R57gen_3: TGTGTGGAAAGTGTGGTGAAGC using N2 genomic DNA as a template. 10 ng/µl of the PCR fragment with 90 ng/µl of marker plasmid gpb-2::GFP (van der Linden et al., 2001) was injected into UF209 gcp-2.1(ok1004) worms. A line transmitting the transgene mosaically, as assessed by GFP expression, was isolated (UF215 gcp-2.1(ok1004) gqEx37[gcp-2.1, Pgbp-2::GFP]). The transgene rescued the phenotypes of the gcp-2.1 mutant.

Image analysis for worm growth. Related to Figure 1

Plates were imaged at 4.0x magnification using a Leica M165 FL stereomicroscope. The images were opened in ImageJ (Schneider et al., 2012) and the zoom function applied so that each image was 150% its original size, to enable more accurate measurement. The freehand line tool was then used to trace along the side of the body of each worm and the resulting line measured. To minimize bias, animals were selected for measurement randomly, using a grid overlay and then a random number generator to specify a grid reference. All animals in this specified square were then measured until a total of 30 worms had been measured for each condition. If an animal occupied two or more squares it was not measured.

Screening method. Related to Figure 4

Temperature sensitive sterile glp-4( bn2) worms were used in the screen. They were maintained at 15°C on OP50 and partly synchronized with an overnight timed egg lay. Three days later they were shifted to 25°C and then on the following day L4 worms were transferred to plates containing the bacterial strains to be
screened (Table S2A). Strains reported by Baba et al. to grow poorly in LB (<0.4 OD after 22 hours) and 15 strains that grew noticeably poorly on NGM plates were excluded (Table S2B). The screen plates were seeded 48 hours beforehand and left to grow at room temperature. The plates contained kanamycin (20 µg/ml) to kill the OP50. OP50-GFP was used to monitor bacterial persistence. For each strain at least 25 worms per plate were placed on at least 3 plates. For each batch, which consisted of 30-80 strains, 9 plates of 25+ worms were set up on a wild type BW21153 transformed with the Kan-containing plasmid pGreen 0029 (WT Kan). Plates were maintained at 25°C and survival was scored at either 11 or 12 days after the first day of adulthood. WT Kan plates were scored at both day 11 and 12. Survival was scored as number of alive worms/total number of worms. Strains that showed at least 15% increased survival (SM) compared to the wild type control (SW) were tested a second round of the screen (Table S3). Strains that passed this round, along with three other strains of interest (Table S3), were verified by PCR and tested in a full lifespan analysis using over 130 worms per strain in which survival was scored every two or three days (Table S3). For strains that increased lifespan significantly in this third round, we complemented the missing gene with a wild type copy carried on a low copy plasmid (Table S3). The mutant ihfA, due to difficulties cloning the complementation construct, was confirmed by testing the “even” strain from the Keio collection (Table S1).

**Estimating batch and plate variation on worm survival. Related to Figure 4A and Figure S2**

First, we quantified the degree of variation in wild type (WT) survival both between batches and between plates within a batch. For simplicity we restricted our analysis to worms that were scored on day 11 (we find similar results when we examine worms scored on day 12). Let B be the number of batches performed and let Pi be the number of WT plates scored on day 11 in batch i. In total, we estimated variation in WT survival by day 11 from 209 plates distributed across B = 30 batches. The number of worms initially on a plate varied from 8 to 37 (mean = 22.66 worms), and the number of plates per batch varied from 3 to 12 (mean = 6.97 plates) (Figure S2Ci). Suppose the variation in the mean WT survival among batches has a beta-distribution with mean \( \bar{\theta} \) and variance \( \bar{\theta}(1 - \bar{\theta})\varphi_b/(1 + \varphi_b) \) (see (Richards, 2008) for details). Also, suppose that variation in mean survival among plates within a batch also has a beta-distribution with variance parameter \( \varphi_p \). These assumptions seem reasonable when inspecting the raw data describing the
between batch and between plate variation in survival fractions (Figure S2Cii). Under these assumptions, if $n_{ij}$ of the $N_{ij}$ worms on plate $j$ in batch $i$ survive to day 11, then the likelihood of the model, described by the set of parameters: $q = \{\theta, \emptyset_b, \emptyset_p\}$, is given by

$$L(\theta|\text{data}) = \prod_{i=1}^{I} f_{p=0}^{1} f_b(p|\theta, \emptyset_b) \prod_{j=1}^{J} f_{bb}(n_{ij}|N_{ij}, p, \emptyset_p) dp,$$

where $f_b$ and $f_{bb}$ are the probability density functions for the beta and the beta-binomial distributions (see (Richards, 2008) for details). Likelihood ratio tests (LRTs) were used to evaluate if $\emptyset_b$ or $\emptyset_p$ were significantly different from zero (i.e. to test if there is no between batch or between plate variation). We found strong evidence of variation in WT survival between plates (LRT, $G_1 = 52.6, P < 0.001$) and variation in survival between plates within a batch (LRT, $G_1 = 11.7, P = 0.001$). Our best-fitting model was described by $q = \{\theta = 0.369, \emptyset_b = 0.042, \emptyset_p = 0.017\}$, which, as expected, predicts between batch variation being greater than between plate variation (Figure S2D).

**Defined medium amino acid mix**

The amino acid mix was designed using the composition of soy peptone. Together with trace metals, the mix was used to replace peptone in NGM (See Experimental Procedures for details). The concentration of the total mix was adjusted downwards so that growth of *E. coli* on defined medium agar plates was similar to that on peptone agar plates. Final concentrations (g/L): Alanine (1.419), Arginine (1.293), Aspartic acid (1.782), Cysteine (0.102), Glutamic acid (2.994), Glycine (3.414), Histidine (0.282), Isoleucine (0.468), Leucine (0.846), Lysine (0.924), Methionine (0.219), Phenylalanine (0.549), Proline (2.064), Serine (0.843), Threonine (0.534), Tryptophan (0.057), Tyrosine (0.318), Valine (0.627).

**Supplemental References**

