Regulated spatial organization and sensitivity of cytosolic protein oxidation in *Caenorhabditis elegans*

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Cells adjust their behaviour in response to redox events by regulating protein activity through the reversible formation of disulfide bridges between cysteine thiols. However, the spatial and temporal control of these modifications remains poorly understood in multicellular organisms. Here we measured the protein thiol–disulfide balance in live *Caenorhabditis elegans* using a genetically encoded redox sensor and found that it is specific to tissues and is patterned spatially within a tissue. Insulin signalling regulates the sensor’s oxidation at both of these levels. Unexpectedly, we found that isogenic individuals exhibit large differences in the sensor’s thiol–disulfide balance. This variation contrasts with the general view that glutathione acts as the main cellular redox buffer. Indeed, our work suggests that glutathione converts small changes in its oxidation level into large changes in its redox potential. We therefore propose that glutathione facilitates the sensitive control of the thiol–disulfide balance of target proteins in response to cellular redox events.
The reversible formation of disulfide bonds between cysteine residues is increasingly recognized as an important mechanism for the regulation of protein function. These post-translational modifications can modulate the activity of a wide range of proteins, including transcription factors, kinases, metabolic enzymes and membrane channels.2–7. As a result, these modifications affect diverse cellular processes, including metabolism, gene expression and cytoskeletal dynamics.7,8. A better understanding of the mechanisms that control these post-translational modifications in vivo may help to explain why the dysregulation of protein oxidation is a common factor in the development of many chronic diseases, including diabetes, cardiovascular disease, neurodegenerative diseases and cancers.9–12. Organisms ranging from bacteria to humans control the formation of protein disulphides in the cytosol through the action of the glutathione and thioredoxin redox systems.13,14. Glutathione (GSH) reduces protein disulphides in a reaction that produces glutathione disulphide (GSSG). Similarly, thioredoxin (trxred) reduces protein disulphides, in a reaction that produces oxidized thioredoxin (trxox). GSSG and trxox are reduced by NADPH in reactions catalysed by specific enzymes.13. As a result, the GSH/GSSG and trxred/trxox couples act as shuttles of electrons between NADPH and protein disulphides.13.

The glutathione and thioredoxin couples have a broad spectrum of distinct but overlapping sets of target proteins.9–12. Both of these couples can affect the formation of disulfides within and between proteins. In addition, the glutathione couple also affects the formation of disulphides between proteins and GSH, which are known to modify the activities of a large number of proteins.9–12. The tendencies of the glutathione and thioredoxin couples to donate electrons to their target proteins are quantified by their redox potentials. Cellular inputs that affect the relative concentrations of these couple’s oxidized and reduced species will shift their redox potential and tilt the thiol–disulfide balance of their respective protein targets. Thus, knowing the redox potential of these couples can inform us about the thiol–disulfide balance of the network of proteins they control.15.

The human and C. elegans proteomes contain ~210,000 cysteine residues, many of which can form disulfides.15,16. The regulation of protein oxidation under the control of the glutathione couple has remained largely unexplored in multicellular organisms due to the limitations of biochemical approaches that generally do not allow to differentiate between cellular compartments, tissues and even individuals.17. The recent development of genetically encoded fluorescent redox sensors that respond to the glutathione couple has enabled studies of the in vivo distribution of this redox potential across subcellular compartments in plants18,19 and across tissues in fruitfly larvae.20. Here we used this approach to visualize the spatial organization of the glutathione redox potential in the cytosol of live C. elegans and quantify its sensitivity and dynamic response. We found that this redox potential is structured at the tissue and sub-tissue levels, and is regulated by insulin signalling at both of these levels. Notably, our work suggests that the glutathione couple is not positioned to act as a buffer in the cytosol, as its redox potential is highly sensitive even to small changes in glutathione oxidation. This sensitivity may enable cells to respond to small perturbations of their cytosolic redox environment by adjusting the thiol–disulfide balance of the network of proteins controlled by the glutathione couple.

Results
Measurement of protein oxidation in vivo. To visualize protein disulphide levels with spatial and temporal resolution in live C. elegans, we used the redox probe roGFP1_R12, reduction-oxidation sensitive green fluorescent protein (roGFP) or ‘sensor’ for short.21. This sensor includes two cysteines whose thiol groups can form a reversible intramolecular disulfide bond. This oxidative modification changes the excitation profile of the sensor’s chromophore by increasing absorption at the 410-nm excitation band and decreasing absorption at the 470-nm band.22. The resulting spectral changes allowed us to monitor the balance between reduced (roGFPred) and oxidized (roGFPox) forms of the sensor via ratiometric fluorescence microscopy.23.

We characterized the response of the sensor in live C. elegans on exposure to exogenous oxidants and reductants that react directly with protein thiols. We treated animals expressing this sensor in the pharyngeal muscles with 50 mM diamide (a thiol-specific oxidant)25 and then 100 mM dithiothreitol (DTT, a reducing agent), and observed the effect of this treatment time course on the sensor’s fluorescence (Fig. 1a–d). Diamide and DTT caused reciprocal changes in fluorescence, indicating that the sensor responds to oxidation and reduction reversibly (Fig. 1c,d). This treatment sequence resulted in the maximal oxidation and reduction of the sensor (see Methods for details). The resulting fluorescence profile during roGFP1_R12 exhibited a large, 7.8-fold dynamic range (Fig. 1b).

Although R410/470 values provide some information about the balance between oxidized and reduced forms of the sensor, they do not represent a biophysically interpretable magnitude. We derived the fraction of sensor molecules with a disulfide bond—OxD1R12, equal to [roGFPox]/([roGFPox] + [roGFPred])—by determining the three conversion parameters relating OxD1R12 and R410/470 from the fluorescence time courses in diamide- and DTT-treated animals (see Fig. 1e legend and Methods for details). The tendency of roGFPox to acquire electrons and thereby become reduced into roGFPred is quantified by the half-cell reduction potential of the roGFPox/roGFPred couple, EroGFPox. This redox potential is given by the Nernst equation EroGFPox = EroGFP - [RT/(2F)] ln ([roGFPred]/[roGFPox]), where R is the gas constant, F the Faraday constant, T the absolute temperature and EroGFP the standard roGFP1_R12 midpoint potential, which is ~ 265 mV (ref. 23). Knowing OxD1R12 made it possible to calculate EroGFP by substituting in the Nernst equation the term [roGFPred]/[roGFPox] with the term (1 – OxD1R12)/OxD1R12, leading to the relation EroGFP = EroGFP - [RT/(2F)] ln [(1 – OxD1R12)/OxD1R12] (Fig. 1e,f).

Previous studies demonstrate that roGFP-based sensors respond to the glutathione couple via glutaredoxin (see Supplementary Note 1). We measured the kinetics of spontaneous recovery from maximal oxidation by transferring animals from diamide treatment back to normal conditions. These in vivo reduction kinetics are fast (Supplementary Fig. 1, t1/2 = 4.1 min), and are comparable to those observed in vitro in the presence of glutaredoxin18,19,21. This suggests that the endogenous levels of glutaredoxin are sufficient to ensure the equilibration of the sensor and glutathione couples in pharyngeal muscle. It is noteworthy that the fraction oxidized of roGFP1_R12 and, consequently, the associated redox potential of the sensor couple are in a steady state in untreated animals (Supplementary Figs 1 and 2) and reach a new steady state close to the original on recovery from diamide (Supplementary Fig. 1). The fast kinetics of reduction of the sensor, together with the observed stability in the sensor’s redox potential over time, suggest that the sensor oxidation is in steady state in unperturbed animals because the redox environment controlling its oxidation is stable.

To determine whether the sensor responds to changes in the total amount of glutathione in live C. elegans, we measured its redox potential in feeding muscles of animals with reduced gcs-1 activity. The gcs-1 gene encodes the only glutamate–cysteine ligase in the genome, which catalyses the rate-limiting first step in
Figure 1 | Measurement of roGFP1_R12 redox potential in live C. elegans. (a) Treatment schedule. Each individual was imaged every minute for 10 min before any treatment and during the last 30 min of the diamide and DTT treatments (grey areas in the top bar). (b–d) Fluorescence intensities and ratios for nine individuals in which the sensor was first maximally oxidized and then maximally reduced by diamide and DTT treatments, respectively. (b) Ratio R_{410/470} of the fluorescence intensities on excitation with 410 nm (c) and 470 nm (d). (e–f) From the R_{410/470} and l_{370} time courses in b and d, respectively, we determined the three parameters required to relate the fluorescence ratio R_{410/470} to the fraction of roGFP1_R12 molecules with a disulfide bond, OxD_{roGFP}. (i) R_{ox}, the value of R_{410/470} when the sensor is fully oxidized; (ii) R_{red}, the value of R_{410/470} when the sensor is fully reduced; (iii) \( \alpha = l_{470} / l_{470+450} \), the ratio of fluorescence intensities on excitation with 470 nm when the sensor is fully oxidized and fully reduced. These three parameters enable the conversion of R_{410/470} to OxD_{roGFP} (ref. 17), shown in e: OxD_{roGFP} = (R_{410/470} – R_{ox})/\alpha = (R_{ox} – R_{410/470}) + (R_{410/470} – R_{ox}). \) The parameters R_{ox} and R_{red} define the dynamic range of the sensor (R_{ox}/R_{red} = 7.8). Next, we used the Nernst equation to convert OxD_{roGFP} into E_{roGFP}, which is the half-cell reduction potential of the sensor redox couple. The parameter \( \alpha \) shifts the relationship from R_{410/470} to E_{roGFP} by a constant value. See Methods for additional considerations. Each of the nine time courses provides one set of three parameters (R_{ox}, R_{red} and \( \alpha \)), yielding a particular OxD_{roGFP} versus R_{410/470} (e) and E_{roGFP} versus R_{410/470} (f) curve. These nine curves are shown in grey. The average value of each parameter (R_{ox} = 5.207, R_{red} = 0.667 and \( \alpha = 0.171 \)) yields the curves shown in black that were used throughout the paper to transform our fluorescent readout R_{410/470} into a biophysically interpretable magnitude E_{roGFP}.

GSH biosynthesis^{26}. Animals with reduced gcs-1 activity exhibit lower GSH levels than wild type^{26}. The gcs-1(ok436) null allele causes larval lethality; therefore, we examined young (L2) larvae lacking zygotic gcs-1, derived from gcs-1 heterozygous parents. We found that the sensor’s redox potential was significantly higher in gcs-1(maternal + zygotic-) animals than in their gcs-1(maternal + zygotic +) siblings (Supplementary Fig. 3). We conclude that roGFP1_R12 responds to changes in GSH synthesis in live C. elegans.

The sensor’s redox potential varies between tissues. To investigate the control of the sensor’s redox potential across tissues of a live animal, we expressed the sensor in the cytosol of cells representing the three primary tissue layers: endoderm (intestine), mesoderm (pharyngeal muscles) and ectoderm (PLM neurons). Unexpectedly, we found that the sensor’s redox potential in each of these tissues varied widely across individuals—up to 13 mV—even though these animals were genetically identical, had the same age and were cultured in the same environment. Despite this individual variation, the sensor’s redox potential differed significantly between these three tissues (Fig. 2 and Supplementary Fig. 4). The intestine had the most oxidizing average redox potential (E_{roGFP} = −268.1 ± 2.1 mV), followed by the PLM neurons (E_{roGFP} = −269.9 ± 1.3 mV) and the pharyngeal muscles (E_{roGFP} = −270.9 ± 1.7 mV). We conclude that the sensor’s redox potential is tissue specific.

The sensor’s redox potential is patterned in the pharynx. To investigate the spatial control of the sensor’s redox potential within a tissue of a live animal, we chose to focus on the pharynx, the feeding organ of C. elegans (Fig. 3a). This large organ has a simple and stereotyped anatomy that facilitates cellular identification. The pharynx is composed of eight adjacent muscles, p1–p8, aligned sequentially and connected by gap junctions^{27}. We acquired profiles of the sensor’s redox potential along the anterior–posterior (A–P) axis of the pharynx for 394...
individuals. We found that these profiles vary considerably across individuals, even though these animals were genetically identical and were cultured in the same environment. At every position along the A–P axis, we observed substantial variation in the sensor’s redox potential between individuals—up to 12 mV (Fig. 3b). Within an individual, the sensor’s redox potentials in pm3, pm4, pm5 and pm7 muscles were strongly correlated (Supplementary Fig. 5). Therefore, the sensor’s redox potential varies in a concerted manner throughout the pharynx.

Surprisingly, we found that the sensor’s redox potential is not uniform throughout the A–P axis of the pharynx but, instead, is spatially patterned (Fig. 3c). We observed three distinct redox regions that align sharply with muscle boundaries (Supplementary Fig. 6a,b and Supplementary Movie 1). These redox regions may encompass multiple muscles, as pm3 and pm4 exhibit no significant difference in the sensor’s redox potential (Supplementary Fig. 6). Along the A–P axis of the pharynx, sensor redox potentials are typically ordered: pm3 = pm4 < pm5 < pm7 (Fig. 3c and Supplementary Fig. 6). Differences between pairs of muscles vary widely between individuals—up to 7.5 mV (Supplementary Fig. 6c,d and Supplementary Movie 2). The redox potential difference between pm3 and pm5 is independent of the difference between pm5 and pm7 (Supplementary Fig. 7), indicating that the redox pattern is generated by two distinct mechanisms. We conclude that the pharyngeal redox profile is patterned by independent mechanisms causing local deviations from the overall glutathione redox potential of the pharynx.

Insulin signalling regulates the sensor’s redox potential. The well-defined redox pattern we observed in the pharynx and the redox-potential differences we observed across tissues suggest that the sensor’s redox potential may be regulated at the tissue and sub-tissue levels. Insulin signalling has evolutionarily conserved effects on survival under oxidative stress8,29, making it a good candidate for the modulation of the sensor’s redox potential. We examined mutants of daf-2, the only insulin receptor gene in C. elegans30, and found that they exhibit a more reducing environment in the pharynx. The daf-2(e1370) mutation, which affects the kinase domain of the protein, and the daf-2(m579) mutation, which affects the ligand-binding domain and is homologous to the human type-A insulin resistance mutation31, lower the sensor’s redox potential by an average of 2.8 mV (Fig. 4a and Supplementary Fig. 8). We observed similar changes in the intestine of daf-2(e1370) mutants (Fig. 4b), but this mutation had no effect on redox potential in the cytosol of PLM neurons (Fig. 4c). Thus, under normal conditions, the effect of daf-2 on redox potential is tissue specific.

Next, we determined whether insulin signalling regulates the spatial redox pattern of the pharynx. We acquired A–P profiles of the sensor’s redox potential in mutants of the FOXO transcription factor DAF-16 whose activity is negatively modulated by signalling through DAF-2 (refs 32,33). We found that daf-16(mu86) null mutants exhibit higher redox potentials in the anterior muscles pm3 and pm5 than wild-type animals (Fig. 4d and Supplementary Fig. 9a). However, this mutation does not affect the sensor’s redox potential in the posterior muscle pm7 (Fig. 4d and Supplementary Fig. 9a). Thus, in wild-type animals, DAF-16 is partially enabled in the anterior muscles of the pharynx and effectively switched off in the posterior. We also found that daf-2(e1370) lowers the sensor’s redox potential at all positions along the pharyngeal A–P axis (Fig. 4d and Supplementary Fig. 9a). This effect is mediated by daf-16, as the redox profiles of the daf-16 single mutants and daf-16; daf-2 double mutants are indistinguishable (Fig. 4d and Supplementary Fig. 9a). We conclude that in contrast to the wild-type case, DAF-16 is enabled in all positions along the A–P axis of the pharynx in daf-2(e1370) mutants. We quantified the relative magnitude of the genetic interaction between daf-2 and daf-16 in the control of the spatial variation of redox potential along the pharyngeal A–P axis using a functional version of categorical regression on genotype34 (Fig. 4e). This analysis indicates that the quantitative regulation of DAF-16 by DAF-2 contributes to the redox pattern of the pharynx.

Glutathione does not act as a redox buffer in the cytosol. As we noted above, the fast kinetics of reduction and the stability of the sensor’s redox potential indicate that this potential equals the redox potential $E_{\text{GSH}}$ of the cytosolic glutathione couple, under unperturbed conditions (Supplementary Figs 1 and 2 and Supplementary Note 1). Glutathione is the most abundant cellular thiol and is widely considered to act as the main redox buffer of the cell13,15,36. Based on this framework, we were surprised to observe a large $E_{\text{GSH}}$ variation between individuals (Figs 2b and 3b), as this variation is likely to cause significant differences in the
the thiol–disulfide balance of proteins targeted by the glutathione system (see sensitivity section below). This prompted us to re-examine the notion that the glutathione couple acts as a redox buffer.

If the glutathione couple acts as a redox buffer, then its redox potential should have a low sensitivity to changes in the balance between reduced and oxidized glutathione species\(^{37}\). We therefore determined whether the redox potentials we measured are located in a region of the Nernst curve where glutathione could be expected to act as a buffer of redox potential. As the oxidation of GSH into GSSG is a bimolecular reaction, \(E_{GSH}^{\text{redox}}\) depends not only on the oxidation state of the glutathione couple (\(Ox^{D}_{\text{GSH}}\)), but also on total glutathione \(GSH_{\text{tot}}\) (\(= [GSH] + 2[GSSG]\)). We placed the \(E_{GSH}^{\text{redox}}\) values we obtained with \(\text{in vivo}\) sensors in the cytosol of cells in the pharynx, intestine and PLM neurons (Figs 2b and 3b) in the context of Nernst curves for different values of \(GSH_{\text{tot}}\). All of these redox potentials fall in a highly sensitive, non-buffering region of the Nernst curve, where even slight changes in \(Ox^{D}_{\text{GSH}}\) lead to large changes in \(E_{GSH}^{\text{redox}}\) (Fig. 5a). For example, in a cell that exhibits the average cytosolic \(E_{GSH}^{\text{redox}}\) of pm3 muscles \((-270 \text{ mV})\), the value of \(Ox^{D}_{\text{GSH}}\) would be 0.0019 assuming a \(GSH_{\text{tot}}\) of 10 mM. The sensitivity of \(E_{GSH}^{\text{redox}}\) to changes in \(Ox^{D}_{\text{GSH}}\) (the slope of the Nernst curve) around this \(Ox^{D}_{\text{GSH}}\) value is 88 times higher than it would be if the glutathione couple were best positioned to buffer redox potential (Fig. 5b). This relative sensitivity rises to 696-fold if we assume that \(GSH_{\text{tot}}\) is 1 mM, at the low end of physiological estimates. We conclude that the glutathione couple does not act as a buffer of redox potential, as previously thought. This result holds for \(GSH_{\text{tot}}\) values up to 1 M.

As an independent confirmation of this finding, we evaluated recent measurements of glutathione oxidation in \(C.\) \textit{elegans} derived from studies that did not involve genetically encoded sensors. These biochemical measurements on whole-worm extracts directly yield \(Ox^{D}_{\text{GSH}}\) values of 0.028 (ref. 39) to 0.035 (ref. 40). These extracts average \(Ox^{D}_{\text{GSH}}\) over all cellular compartments, including the highly oxidizing endoplasmic reticulum, where \(Ox^{D}_{\text{GSH}}\) equals 0.4 (ref. 41). It is not surprising, therefore, that these reported values are higher by an order of magnitude than the \(\text{in vivo}\) values we obtained with
Regression on genotype. This regression expresses the A–P redox profile in terms of the wild-type profile plus effect terms derived from redox potential along the A–P axis of the pharynx. We quantify the effect size of (16(mu86) (red, 224 animals) and the double-mutant panel shows that in most of the pharynx of wild-type animals curves divided by the difference between the blue and the orange curves. The latter difference is the maximal effect of single and double mutants (Supplementary Fig. 9b,c). In essence, the normalized effect of signalling in the cytosolic redox potential. Cumulative distributions of cytosolic redox potentials in wild type and in total glutathione were the main source of redox potential determination whether the magnitude of the animal’s redox response on shift to a more oxidative environment is predicted by the redox potential of the glutathione couple before such shift. If differences in total glutathione were the main source of redox potential variation across individuals, the sensitivities associated with different redox potentials would exhibit little or no variation and, therefore, we would not observe a correlation between redox potential and redox displacements on oxidative shift. On the other hand, if the variation in redox potential across individuals stemmed from differences in the fraction of oxidized glutathione, we would observe a negative correlation between redox potential and the change in redox potential on oxidative shift. In the absence of this Nernst framework, the latter case appears counter intuitive, as it states that animals with higher levels of protein oxidation before oxidant exposure would be less sensitive to oxidant treatment than animals with lower initial oxidation levels.

Glutathione oxidation varies between individuals. What are the mechanisms that cause isogenic animals to have different sensor’s redox potentials? The observed differences in redox potential between individuals could be explained by individual differences in the fraction of oxidized glutathione. However, they could also be due to individual differences in total glutathione content. The properties of the Nernst curve allow us to determine which of these two scenarios is dominant, because changes in glutathione oxidation state affect sensitivity (the slope of the Nernst curve, Fig. 5b), whereas changes in total glutathione do not affect sensitivity at any glutathione oxidation state (they only shift the Nernst curve vertically, Fig. 5a). To distinguish between these possibilities, we determined whether the magnitude of the animal’s redox response on shift to a more oxidative environment is predicted by the redox potential of the glutathione couple before such shift. If differences in total glutathione were the main source of redox potential variation across individuals, the sensitivities associated with different redox potentials would exhibit little or no variation and, therefore, we would not observe a correlation between redox potential and redox displacements on oxidative shift.
sensor’s redox potential of pm3, pm5 and pm7 muscles relative to their baseline redox potential (defined as the average redox potential before oxidant exposure; Fig. 6b–d and Supplementary Fig. 10). Each of these muscles exhibits a distinct response (Fig. 6e). Although all muscles experience an increase in the sensor’s redox potential after oxidant exposure, only pm5 and pm7 exhibit an adaptive response consisting of a rise and subsequent decrease in redox potential. In addition to these regularities in the response behaviour of each muscle, we observed a substantial variation across individuals in the magnitude of the response to oxidant treatment (Fig. 6b–d, and Supplementary Figs 10 and 11). Animals with a higher baseline (coloured in red, Fig. 6b–d and Supplementary Fig. 10) exhibit smaller redox displacements than animals with a lower baseline (coloured in blue). This relation is confirmed by the negative sign of the observed strong correlations between baselines and early redox displacements of pm3, pm5 and pm7 muscles (Fig. 6f). Thus, the response dynamics that we observe indicate that differences in baseline across individuals are due in large part to differences in glutathione oxidation, not glutathione content.

Sensitivity of protein oxidation to $E_{GSH}$ variation. In our study we encountered differences in the sensor’s redox potential across individuals (13 mV), across tissues (2.8 mV), within tissues (7.5 mV), across genetic perturbations (4.2 mV) and under oxidative stress (15 mV). To understand how much these differences may impact protein oxidation, we considered the effect of $E_{GSH}$ variation on the thiol–disulfide balance on protein targets in equilibrium with the glutathione couple.

Changes in the glutathione redox potential sway the oxidation states of target proteins in the same direction, but the magnitude depends on both the protein’s tendency to remain reduced (quantified by its midpoint potential) and its initial oxidation state (determined by the redox potential it was experiencing). Midpoint potentials are presently unknown for most proteins. Still, we can treat them as variables and analyse the impact of a redox-potential change on protein oxidation, both in terms of absolute differences and fold changes (Fig. 7). For example, a difference in glutathione redox potential of 10 mV that we typically observe in response to 5 mM t-BuOOH treatment would affect the oxidized fraction of a protein by at most 19% (absolute difference; Fig. 7b) or by a factor of at most 2.2 (fold change; Fig. 7c). Such changes can be biologically significant, especially given the wide variety of redox-sensitive proteins.

Discussion

The study of the regulation of cytosolic glutathione redox potential has been challenging, especially in live multicellular organisms. The methods used until recently required the homogenization of samples, resulting in the mixing of subcellular compartments, cells, tissues and often entire populations. In this study, we overcame this difficulty by using a genetically encoded protein sensor. The ability to visualize and quantify the sensor’s redox potential in live individuals, at cellular and high temporal resolution, made it possible to investigate the structure and regulation of the cytosolic glutathione redox potential, as well as the determinants of its variation. As we discuss below, this variation reflects, in part, the high sensitivity of this potential to changes in glutathione oxidation. The regulation of the sensor’s redox potential is suggested by (i) differences in the average redox potentials of the pharynx, intestine and PLM neurons; (ii) the spatial patterning of the sensor’s redox potential along the A–P axis of the pharynx; and (iii) differences in the response dynamics to oxidant treatment of individual pharyngeal muscles.

Figure 5 | The cytosolic glutathione couple is not a buffer of redox potential. (a) The redox potential of glutathione thiol (GSH)/glutathione disulfide (GSSG) couple is given by the Nernst equation $E_{GSH} = E_{GSH}^- \cdot \left(\frac{RT}{2F}\right) \ln \left(\frac{[GSSG]}{[GSH]}\right)$, where $R$ is the gas constant, $F$ is the Faraday constant and $T$ is the absolute temperature, here 295.15 K. The midpoint (standard) potential $E^*$ for the glutathione couple is $-240$ mV under biological standard conditions. This equation can be rewritten in terms of total glutathione $GSH_{tot}$ (equal to 2 [GSSG] + [GSH]) and the fraction of oxidized glutathione $OxD_{GSH}$ (equal to 2 [GSSG]/$GSH_{tot}$), resulting in the expression $E_{GSH} = E_{GSH}^- \cdot \left(\frac{RT}{2F}\right) \ln \left(2 \cdot [GSSG]/GSH_{tot} \cdot (1 - OxD_{GSH})^2/OxD_{GSH} \right)$. The panel shows the Nernst curves for $GSH_{tot} = 1, 10$ and 20 mM. A value of 10 mM is at the high end of physiological concentrations observed in other species. Changes in the oxidation state of the glutathione couple will cause a large change in redox potential when the couple is in a highly sensitive (red) region of the curve and will cause a small change in redox potential when the couple is in a buffered (green) region of the curve. The coloured circles indicate the intersection of the average cytosolic redox potentials we measured in the pharyngeal muscles, intestine and PLM touch neurons (Fig. 2) with the 10-mM $GSH_{tot}$ curve. (b) Normalized sensitivity of the redox potential $E_{GSH}$ to changes in $OxD_{GSH}$ (defined as the derivative of the Nernst curves shown in a, normalized to its minimum value). $E_{GSH}$ sensitivity is independent of $GSH_{tot}$. Normalized sensitivity values below 1.5 indicate that the glutathione couple operates in a buffering regime (0.157 < $OxD_{GSH}$ < 0.729, green), while values above 5 indicate a highly sensitive regime ($OxD_{GSH}$ < 0.037, red; $OxD_{GSH}$ > 0.929, not highlighted). The glutathione couple operates in a highly sensitive regime in vivo (for example, at $OxD_{GSH}$ = 0.0019 associated with $E = -270$ mV, black arrow).
Figure 6 | Variation in glutathione redox potential between individuals is caused by differences in glutathione oxidation. (a) Spatial and temporal response of the sensor’s redox potential in an individual before and during treatment with 5 mM t-BuOOH. Each column represents the redox profile of the individual along the A–P pharyngeal axis at a given time point. Each row is a time point at a given spatial location. Worms were imaged every 30 s. Oxidant exposure starts at $t = 0$ (grey background in b–f). (b–d) The spatiotemporal response to 5 mM t-BuOOH treatment was recorded in 64 individuals. For each individual, we determined the responses of pm3, pm5 and pm7 (b,c and d, respectively). Each time series was baseline corrected by subtracting its average potential before oxidant exposure. The resulting time series were converted into a continuous function using a spline basis (for the non-baseline corrected set see Supplementary Fig. 10; for the quality of the fit see Supplementary Fig. 11). The colouring encodes the average potential before oxidant exposure ($E_b$). (e) Average response dynamics. The functionalized trajectories of redox potential in pm3 (blue), pm5 (green) and pm7 (red) were pooled and subjected to functional categorical regression\(^3\) on the regions of origin (pm3, pm5 and pm7). This procedure yields the average time-dependent response dynamics in each region. Shaded areas represent 68% point-wise confidence intervals. (f) For each muscle, the Pearson correlation between the individual redox baselines before perturbation and the baseline-corrected response after perturbation. The strong initial negative correlation implies that we mostly see different baseline potentials on the same Nernst curve rather than different curves corresponding to distinct values of total glutathione (see text for details). The subsequent correlation decay indicates that an adaptive response to perturbation is becoming prominent. Shaded areas represent 68% point-wise confidence intervals.

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We found that the action of insulin signalling affects the spatial organization of the sensor’s redox potential. The insulin receptor DAF-2 regulates this redox potential in the cytosol in a tissue-specific manner, affecting it in pharyngeal muscles and intestine, but not in PLM neurons. In addition, DAF-2 contributes quantitatively to the spatial patterning of redox potential in the pharynx by regulating the activity of the FOXO transcription factor DAF-16. We also found that a daf-2 mutation that causes type-A insulin resistance in humans is sufficient to lower the cytosolic redox potential of the glutathione couple, which may lead to a significant decrease in the oxidation of target proteins. It would be interesting to investigate whether similar changes in the thiol–disulfide balance are important to the pathogenesis of insulin resistance and diabetes in humans.

Glutathione has been widely thought to act as the main redox buffer of the cell\textsuperscript{1-3}, a function that provides homeostasis by protecting proteins from the indirect oxidizing action of reactive oxygen species. This view seemed difficult to reconcile with the considerable variation in glutathione redox potential across isogenic individuals, as this variation might cause significant differences in protein oxidation levels. In an effort to substantiate whether the glutathione couple acts as a redox buffer in the cytosol, we found that it does the opposite.

The notion of glutathione as a redox buffer may have gained traction over the last half century, because the abundant GSH restores protein thiols that underwent oxidative attack and reduces oxidants such as hydrogen peroxide. Yet, this very action produces an oxidant, GSSG. As most cytosolic glutathione is reduced in vivo, an ever so slight change in GSSG concentration strongly affects the glutathione redox potential (Fig. 5)\textsuperscript{12}. Changes in the glutathione redox potential will have repercussions on a much wider array of cysteine thiols than those that were initially threatened by oxidative attack, because cysteine thiols generally react in vivo very slowly with hydrogen peroxide but much faster with GSSG\textsuperscript{12,44}.

Our results call for a change in perspective on the role of the cytosolic glutathione couple. Numerous redox couples, including H$_2$O$_2$/H$_2$O and NADP$^+$/NADPH, interact with the glutathione couple. As a result, the relative levels of GSH and GSSG are determined by a wide range of concurrent processes, including those that generate hydrogen peroxide and those that influence the production or utilization of NADP$^+$ and NADPH. As the glutathione couple is not positioned to buffer its redox potential in the cytosol, this potential responds sensitively to changes in glutathione oxidation. We therefore propose that the glutathione couple integrates and amplifies the effects of diverse redox inputs into a single physiological signal: the glutathione redox potential (Fig. 8), to which proteins with embedded cysteines can respond differentially.

The glutathione couple can be thought of as a broker that mediates the indirect effects of oxidants and reductants on the thiol–disulfide balance of many proteins. As the activity of proteins can be affected by this balance, the high sensitivity of $E_{\text{GSH}}$ to changes in glutathione oxidation may enable cells to respond to very small changes in the concentration of these

Figure 7 | Analysis of the sensitivity of protein oxidation to changes in glutathione redox potential. (a) Relationship between redox potential and protein oxidation. The Nernst equation relates the redox potential $E$ to the fraction oxidized OxD of a redox couple. This relation depends on the couple’s midpoint potential $E^\circ$ (which quantifies the affinity for electrons of the oxidized species). The panel depicts the Nernst curve relative to the midpoint $E^\circ$, that is, OxD versus $E - E^\circ$, with OxD = $1$/($\exp[(-E - E^\circ)/RT] + 1$) and $T = 295.15 \, ^\circ\text{C}$. See Supplementary Note 2 for additional considerations. (b) Effect of a redox-potential change on absolute protein oxidation levels. The calculation of the change in OxD on a change in potential from $E_i - E^\circ$ to $E_f - E^\circ$ (that is, $\Delta E = E_f - E_i$), depends on whether the relevant biochemical quantity is the ensuing difference OxD$_f$ – OxD$_i$ (this panel) or fold-change OxD$_f$/OxD$_i$ (c). The abscissa shows the initial redox potential $E_i$ of a protein relative to its midpoint $E^\circ$. The potential $E_i$ entails a certain fraction of oxidized protein OxD$_i$. When the potential changes from $E_i$ to $E_f$, the fraction of protein oxidized changes accordingly from OxD$_i$ to OxD$_f$. This change is reported by the various curves, one for each potential difference $\Delta E = E_f - E_i$, shown in the legend. (c) Effect of a redox-potential change on relative protein oxidation levels. Similar to (b), but the change of interest is now the ratio of OxD$_f$ at $E_f$ to OxD$_i$ at $E_i$. Note that this is a log-linear plot. The largest fold-change occurs when the initial OxD is as small as possible, as then even the smallest change in OxD results in a dramatic fold change. This occurs at very negative $E_i$ values, far to the left of the midpoint. The fold change will be very large (very small) for positive (negative) $\Delta E$. If the protein couple is far to the right of the midpoint, OxD is near maximal and only small changes can occur, resulting in a fold change close to 1.
reactive chemical species, well before their concentration is high enough to be toxic. This form of intracellular signalling may enable cells to mount an adaptive response to counteract the toxicity of a subsequent and more massive exposure to one of these reactive species. Indeed, disulfide bond formation in direct regulators of Nrf2 and NFκB increases transcription by these factors, which plays an important role in the cellular response to oxidants.

What would happen if the glutathione couple acted as a redox buffer? We expect that thiol–disulfide modification of in principle responsive proteins would become insensitive to redox events that affect the fraction of oxidized glutathione, as those events would no longer entail significant changes in redox potential. For example, the same change in glutathione oxidation that results in a 10 mV increase in $E_{\text{GSH}}$ when the couple is in a highly sensitive region of the Nernst curve ($Ox_{\text{GSH}} = 0.0019$, $GSH_{\text{tot}} = 10$ mM) would only cause a 0.165 mV increase if the couple were optimally positioned as a buffer. As can be seen in Fig. 7, under buffering conditions, such an event would no longer lead to a change in the thiol–disulfide balance of the protein network targeted by glutathione.

As the glutathione couple is so abundant, the concentration of GSSG in the cytosol remains significant compared with the typical concentrations of cytosolic proteins, even though most glutathione is in the reduced form. We estimate that GSSG concentration is about 9 µM in the average pm3 muscle of *C. elegans* ($Ox_{\text{GSH}} = 0.0019$, assuming $GSH_{\text{tot}} = 10$ mM). This concentration is comparable to the 1 µM median concentration of cytosolic proteins in yeast and is only slightly lower than the concentration of abundant cytosolic proteins (for example, the concentrations of ten glycolytic enzymes in mammalian muscle ranges between 29 and 131 µM). A 9 µM GSSG concentration is also much higher than the 0.001–0.7 µM physiological concentration range of cytosolic hydrogen peroxide in aerobic organisms. Therefore, the kinetics of thiol–disulfide exchange between the glutathione couple and its target protein couples are not likely to be limited by the concentrations of GSSG and GSH. This means that the glutathione couple is well positioned to mediate efficiently the transfer of electrons to and from its target proteins.

The differences in glutathione redox potential that we observe in the *C. elegans* pharynx reflect the population’s diversity rather than the individual’s variation over time, as individuals have essentially stable glutathione redox potentials over the course of almost an hour (Supplementary Fig. 2). It is notable that individuals can exhibit distinct and stable glutathione redox potentials, as those differences must arise as a result of minuscule differences in glutathione oxidation. For example, the entire 12 mV range of pharyngeal $E_{\text{GSH}}$ values would be due to $Ox_{\text{GSH}}$ values between 0.0012 and 0.0028, assuming $GSH_{\text{tot}} = 10$ mM for all animals. This means that *C. elegans* is able to control glutathione oxidation with great precision. Thus, the thiol–disulfide balance of the network of protein targets of the glutathione couple is capable of both stability (as a result of the control of glutathione oxidation) and sensitivity (as a result of the high-sensitivity of glutathione redox potential to changes in glutathione oxidation).

The vast majority of cytosolic glutathione (> 99%) is found in the reduced state in a wide range of cell types and organisms. For example, cytosolic Ox$_{\text{GSH}}$ is 0.00004 in yeast, 0.0007 in cultured human HeLa cells, 0.0066 in rat liver and 0.009 in *Escherichia coli*. When placed in the context of the Nernst curve, these Ox$_{\text{GSH}}$ values indicate that the glutathione couple is positioned in the highly sensitive, non-buffering region of the curve in all of these organisms (Fig. 5). Thus, this property is not unique to *C. elegans*.

In the *C. elegans* pharynx, variation in Ox$_{\text{GSH}}$ values between individuals leads to differences in the magnitude of their response to t-BuOOH treatment. Similarly, variation in Ox$_{\text{GSH}}$ values across species suggests large differences in redox sensitivity to changes in glutathione oxidation. For example, we predict yeast to be 159-fold more sensitive than rat liver cells to a small change in Ox$_{\text{GSH}}$. It would be interesting to explore what adaptive constrains caused species to position their glutathione couple to provide different redox sensitivities.

Although the glutathione couple facilitates a sensitive regulation of the thiol–disulfide balance of target proteins in the cytosol, it may well play a buffering role in other subcellular compartments. Redox potential is optimally buffered by the glutathione couple when Ox$_{\text{GSH}}$ equals 0.414 (Fig. 5b). This optimal value is in excellent agreement with measured Ox$_{\text{GSH}}$ values of 0.4 in the endoplasmic reticulum. We conclude that unlike in the cytosol, the glutathione couple is positioned to buffer the thiol–disulfide balance of target proteins in the endoplasmic reticulum.

The glutathione couple’s capacity to sensitively affect the thiol–disulfide balance of its target protein network may be advantageous by enabling organisms to respond to small perturbations to the cytosolic redox environment. However, this sensitivity could also amplify the effects of any breakdown in cellular redox homeostasis, setting the stage for the association between cytosolic redox changes and the aetiology and progression of many human diseases, particularly those for which ageing is a strong risk factor.

Methods

**Strains and culture.** *C. elegans* were cultured under standard conditions at 20 °C. Wild-type *C. elegans* was Bristol N2.

**Construction of transgenes.** We built roGFp1_R12 from the vector pPD96_32 (Fire vector kit, Addgene), which contains a green fluorescent protein (GFP) construct optimized for expression efficiency in *C. elegans*. We replaced seven residues in this construct (C48S, C65S, S147C, N149K, S202K, Q204C and F223R)
by site-directed mutagenesis (QuickChange, Stratagene) to generate roGFP1_R12 (ref. 23). We used a combination of classical cloning and fusion PCR to remove the mitochondrial-targeting sequence in the original plasmid and to fuse the roGFP1_R12 coding sequence to specific promoters. The fusion PCR for the coding region was done with 12 cycles of amplification to minimize the chances of introducing sequence errors (Expand High Fidelity PCR System, Roche), and cloned into a TOPO BluntII vector. We confirmed the sequence for all the constructs injected by DNA sequencing, using the primers listed in Supplementary Table 4.

roGFP1_R12 expression in pharyngeal muscle. We amplified the myo-2 promoter from pPD00.08 (Fire vector kit, Addgene) with primers WF45 and WF46, and cloned the PCR product into a TOPO BluntII vector. roGFP1_R12 was inserted into this vector backbone by restriction with AgeI and BstWI, generating Pmyo-2-roGFP1_R12.

roGFP1_R12 expression in intestine. We amplified the sur-5 promoter from pPD158.87 (Fire vector kit, Addgene) with primers WF45 and WF49, and cloned the PCR product into a TOPO BluntII vector. We fused this promoter to a restriction fragment containing roGFP1_R12 by secondary PCR using primers WF38 and WF129, generating Psur-5-roGFP1_R12, which was cloned into a TOPO BluntII vector. Although we expected the sur-5 promoter to drive roGFP1_R12 expression in several tissues, we observed high expression level for roGFP1_R12 only in the intestine at the ×10 magnification we used for imaging pharyngeal muscles throughout this study.

Generation of transgenic animals and construction of strains. The plasmids containing roGFP1_R12 and Pmyo-2-roGFP1_R12 were isolated using midiprep (Amresco) as coinjection marker. The plasmid containing Pmec-4::roGFP1_R12 was microinjected without a coinjection marker. Pmyo-2-roGFP1_R12, Pmec-4::roGFP1_R12 and Psur-5::roGFP1_R12 were microinjected into wild type at centromers ranging from 15 to 50 µg ml⁻¹, to generate ydEx20 and ydEx42, ydEx37 and ydEx42, and ydEx25, ydEx22 and ydEx42, respectively. ydEx20 was integrated into the genome with ultraviolet light (254 nm, generation yf6). After integration, the strain was outcrossed six times to wild type. Double and triple mutants were generated by standard genetic methods. Unless noted, measurements in pharynx, PLM neurons and intestine were performed in ydIs1, ydEx37 and ydEx25 animals, respectively, gc-1(ok436)/unc-119(e165) J. We performed live fluorescence measurements in Petri dishes with 4 µl 4 × 4 binning using an average exposure time of 40 ms (Figs 1, 2, and 4). Higher spatial resolution images were acquired with 2 × 2 binning, with average exposure times of 165 ms and 300 ms, respectively for Fig. 3 and Supplementary Fig. 3. In control experiments, we imaged 20 animals sequentially with 2 × 2 and 4 × 4 binning, and found that the resulting estimates of the ratio of fluorescence (R410/470) are highly correlated (r = 0.961, P < 0.0001). To image the intestine, we concentrated on the posterior region of the tissue, which exhibits higher sensor expression. Fluorescence images were acquired with 4 × 4 binning, using average exposure times of 105 ms. To image mechanosensory neurons, we selected the PLM neurons because they are located at the tail of the worm—therefore, the IMF was thinner and flat and, therefore, more suitable for imaging. Of all the PLM neuronal images of the 410 nm wavelength, 30 animals were quantified for each worm. Fluorescence images were acquired with 2 × 2 binning using average exposure times of 450 ms.

Optimization of microscopy for different tissues. We optimized exposure times to use at least two thirds of the dynamic range of the camera. This step ensures that image segmentation is performed consistently across all experimental conditions and with all imaged tissues, making it an automated imaging pipeline. The camera exposure times were adjusted to ensure the quality of the alignment to the midline of each pharynx. We refer to this as an ‘alignment’ process for the researcher. We chose the camera binning based on the spatial resolution required for tissues of different morphology and size. To image the feeding muscles of the pharynx, we focused on the plane of the pharyngeal lumen under transmitted light. Fluorescence images were acquired with 4 × 4 binning using an average exposure time of 40 ms (Figs 1, 2, 4 and 6). Higher spatial resolution images were acquired with 2 × 2 binning, with average exposure times of 165 ms and 300 ms, respectively for Fig. 3 and Supplementary Fig. 3. In control experiments, we imaged 20 animals sequentially with 2 × 2 and 4 × 4 binning, and found that the resulting estimates of the ratio of fluorescence (R410/470) are highly correlated (r = 0.961, P < 0.0001). To image the intestine, we concentrated on the posterior region of the tissue, which exhibits higher sensor expression. Fluorescence images were acquired with 4 × 4 binning, using average exposure times of 105 ms. To image mechanosensory neurons, we selected the PLM neurons because they are located at the tail of the worm—therefore, the IMF was thinner and flat and, therefore, more suitable for imaging. Of all the PLM neuronal images of the 410 nm wavelength, 30 animals were quantified for each worm. Fluorescence images were acquired with 2 × 2 binning using average exposure times of 450 ms.

Image processing and segmentation. Image processing was conducted in ImageJ (NIH) and Matlab (Mathworks). We performed background subtraction by removing the mode intensity value of the entire image from each pixel. This procedure removes the background due to the agar and the camera noise, as the vast majority of pixels in our images were part of the background. To segment the images, we defined regions of interest (ROI) by applying a threshold in the 410 nm excitation images. We used the same intensity value for the segmentation across all tissues and conditions. In the case of the PLM neurons, the ROIs correspond to single cells; in the case of the intestine, the ROIs correspond to the most posterior portion of the tissue (~85 µm in length); in the case of feeding muscles, the ROIs correspond to the entire pharynx. We used the ROIs to quantify both the 410 and 470 nm excitation images (I410 and I470, images, respectively). We obtained nearly identical R410/470 values by segmenting based on I410 or I470 images (r = 0.999, P < 0.0001, for the individuals in Fig. 3).

Ratiometric measurements. We calculated R410/470 by computing the ratio of the total fluorescence between the segmented I410 and I470 regions. To study how R410/470 and its transforms vary along the A-P axis of the pharynx, we generated R410/470 profiles along the midline of this tissue using automated scripts in ImageJ. First, the segmented I410 and I470 regions were rotated, centered and referred to orient them along the A-P axis. Then, we constructed a polyline in the I410 image along the midline of the pharynx using features of five morphological pharyngeal landmarks. This polyline was visually inspected and, when necessary, adjusted manually to ensure the quality of the alignment to the midline of each pharynx. We refer to this polyline as the ‘medial axis of the pharynx’, and measurements conducted along the medial axis as ‘anterior–posterior pharyngeal profiles’. We measured the intensity along the medial axis in I410 and I470 images, using a line width of 5.16 µm (2 pixels) for 4 × 4 binning images and using a line width of 7.74 µm (6 pixels) for 2 × 2 binning images. The resulting intensity vectors were then length normalized using bilinear interpolation. We determined the E value profile along the A-P axis of the pharynx from the computed element-wise intensity ratios (I410/I470) between these intensity vectors. We note that the E value (ranging from 0 to 1, computed from the ratio of 410 to 470 total fluorescence along the medial axis) approximates very closely that of the whole pharynx (r = 0.9988, P < 0.0001 for the 394 animals in Fig. 3). To calculate the value of E for individual muscle segments, we identified muscle boundaries based on the position of the morphological landmarks of the pharynx in the length-normalized intensity vectors. We computed E from the ratio of the 410 and 470 nm total intensities within each region.

Image visualization. Pixel-by-pixel E images, computed from the raw unsegmented I410 and I470 images, provide a valuable visualization tool that complements the formal image analysis and quantification described above. We computed E from pixel-by-pixel R410/470 values and generated red, green, and blue (RGB) images containing E values that provide insight into the spatial distribution of sensor expression in the system. We then provided information about the magnitude of the intensity signals used, de-emphasizing pixels where the signal was low. To this end, we adjusted the brightness of pixels...
which intensity values in the L30 and L70 images were below the value of the intensity cutoff used for segmentation (equal to 2,000 in all images shown). Images were transformed using the saturation-brightness colour space and pixel intensities were converted to the hue-saturation-brightness colour space and pixels with intensity values above the cutoff were assigned a brightness value equal to their intensity divided by the cutoff value. The resulting hue-saturation-brightness image was transformed to RGB 0–255 for visualization. We used the cool–warm and similar diverging “colormaps” to colour-code images and pharyngal E profiles, as these colormaps are perceptually linear and behave well for observers with colour-deficient vision.

Determination of OxD$_{GFPR}$ and E$_{GFPR}$ from R$_{410}/470$. To calculate the fraction of roGFPR1_R12 molecules with a disulfide bond (OxD$_{GFPR}$) from ratiometric fluorescence measurements, we determined empirically the value of the three parameters relating the roGFPR1_R12 half-cell redox potential (E$_{GFPR}^R$) as described in the legend of Fig. 1e. We note that the parameter $\sigma$ determines the curvature of the relationship from R$_{410}$/R$_{470}$ to OxD$_{GFPR}$ (if $\sigma$ were equal to 1, then the relationship in Fig. 1e would be linear). We treated worms with various concentrations of oxidants and reductants, and found that the sequential treatment with the oxidant diamide (50 mM, Sigma) followed by treatment with the reducing agent DTT (100 mM, Sigma), resulted in maximal oxidation and reduction of the sensor, respectively, in the pm3 muscles of the pharynx. The in vivo dynamic range of the sensor matched the reported in vitro dynamic range. These experiments also showed that 50 mM diamide treatment and 100 mM DTT treatment affect the expression of the roGFPR1_R12 sensor and GFP S65C by at most 10%, on the timescale of the Fig. 1 experiment. All chemical incubations were performed in NIM plates. No photobleaching was observed during incubations.

We calculated the potential of the redox couple defined by oxidized and reduced forms of roGFPR1_R12 using the Nernst equation $E_{GFPR}^R = E_{GFPR}^R - RT/2F \ln [1-(OxD_{GFPR}/E_{GFPR}^R)]$ and the standard redox potential of roGFPR1_R12 (E$_{GFPR}^R$) of ~265 mV$^{23}$, a temperature of 295.15 K (22 °C) and assuming a cytosolic pH of 7. The final relation from R$_{410}$/R$_{470}$ to E$_{GFPR}$ is given by the expression $E_{GFPR} = E_{GFPR}^R - RT/2F \ln [1-(OxD_{GFPR}/E_{GFPR}^R)]$. It is noteworthy that the parameter $\sigma$ shifts E$_{GFPR}$ by a constant value equal to ~$-RT/2F$ in ln (x). As a result, E$_{GFPR}$ differences are not affected by this parameter. We also note that although the value of E$_{GFPR}$ is influenced by pH (by shifting the midpoint potential of the couple), under physiological conditions these effects are not relevant to understand the equilibration of potentials between roGFPR1_R12 and glutathione couples$^{1,17}$. This is because both couples involve the exchange of two protons and two electrons and, under physiological conditions, pH is well below the pK$_{a}$ of the reactant thiols (pK$_{a}$ = 8.92 for GSH and ~9.0 for roGFPR1_R12 C147 and C204); as a result, the midpoint potentials of these redox couples is ~$-255$ mV$^{23}$. Therefore, roGFPR1_R12 is a pH-insensitive redox sensor.

Functional data analysis. We used the functional data analysis statistical package in MatLab$^{26}$ to model and analyse position series of E-values in pharyngeal A–profiles and time series of E-values in the response to t-BuOOH. The basic idea of this approach is to express discrete spatial or temporal series of observations in the form of a function. These functions are then treated as single observations, which are analysed statistically applying concepts of multivariate analysis$^{24}$. A more detailed description of our application of this technique is given in the Supplementary Methods.

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Author contributions

C.R.-A. and J.A. conceived, designed and performed experiments, and analysed data. C.R.-A. and J.A. provided guidance. C.R.-A., J.A. and C.R.-A. generated roGFP1_R12 vector. J.A. and W.F. provided guidance. C.R.-A. and J.A. generated roGFP1_R12 vector and performed experiments and analysed data. C.R.-A. and J.A. conceived, designed and performed experiments, and analysed data. C.R.-A. and J.A. generated roGFP1_R12 vector. J.A. and W.F. provided guidance. C.R.-A. and J.A. conceived, designed and performed experiments, and analysed data.

Additional information

Supplementary Information accompanies this paper at http://www.nature.com/naturecommunications

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