1-Undecene from *P. aeruginosa* is an olfactory signal for flight or fight response in *C. elegans*

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**Review Timeline:**

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**Transaction Report:**

(Note: With the exception of the correction of typographical or spelling errors that could be a source of ambiguity, letters and reports are not edited. Depending on transfer agreements, referee reports obtained elsewhere may or may not be included in this compilation. Referee reports are anonymous unless the Referee chooses to sign their reports.)
Dear Dr. Singh,

Thank you for submitting your manuscript to The EMBO Journal. Your study has now been seen by three referees and their comments are provided below.

As you can see from the comments, the referees find the analysis interesting. However significant revisions are also needed in order to consider publication here. Important controls are missing, some of the data needs to be better substantiated and further insight into how the induction of the host immune response pathway is linked to survival is needed. Should you be able to extend the findings along the lines suggested by the referees then I would be willing to consider a revised version.

I am happy to discuss the raised points further and maybe it would be most helpful to do so via phone or video.

When preparing your letter of response to the referees' comments, please bear in mind that this will form part of the Review Process File, and will therefore be available online to the community. For more details on our Transparent Editorial Process, please visit our website: https://www.embopress.org/page/journal/14602075/authorguide#transparentprocess

We generally allow three months as standard revision time. As a matter of policy, competing manuscripts published during this period will not negatively impact on our assessment of the conceptual advance presented by your study. However, we request that you contact the editor as soon as possible upon publication of any related work, to discuss how to proceed. Should you foresee a problem in meeting this three-month deadline, please let us know in advance and we may be able to grant an extension.

Thank you for the opportunity to consider your work for publication. I look forward to discussing your revisions further with you

Yours sincerely,

Karin Dumstrei, PhD
Senior Editor
The EMBO Journal

Instructions for preparing your revised manuscript:

Please make sure you upload a letter of response to the referees' comments together with the revised manuscript.

Please also check that the title and abstract of the manuscript are brief, yet explicit, even to non-specialists.

When assembling figures, please refer to our figure preparation guideline in order to ensure proper formatting and readability in print as well as on screen:
https://bit.ly/EMBOPressFigurePreparationGuideline
In this manuscript, the authors report that a Pseudomonas specific volatile odor elicits behavioral avoidance and induces immune-specific gene expression, increasing survival in response to pathogenic bacteria. There is evidence that host immune responses may be driven by chemosensory cues in multiple systems, although the molecular mechanisms of these responses are largely unknown. Recently, it was demonstrated that another C. elegans olfactory neuron - AWC - plays a critical role in non-autonomous regulation of p38 MAPK dependent immune responses via unidentified odors (Foster et al., 2020). Thus, the identification of a parallel olfactory input, including the identity of both the odors and molecular pathways involved, would be a really novel finding of broad interest.

The authors show that 1-undecene elicits behavioral avoidance and induces immune response genes and that undecene production by Pseudomonas is necessary for these responses. This is a really interesting finding, however, at this point there are many cases in which missing controls and/or an insufficiently fleshed out pathway severely limits the interpretation of these data. Additionally, it is unclear how the innate immune responses shown here are coordinated with the observed acute behavioral responses to alter survival.
Specific comments:

1. A major concern is the lack of both genetic and chemical controls for most of the experiments. For all of the C. elegans mutants, only a single allele was analyzed and no rescue experiments were performed. This is important to not only control for genetic background, but also to confirm the specificity of the proposed AWB neuron function. For example, odr-3 is expressed in AWB and AWC, as noted by the authors, but also in ASH, where it is necessary for odor responses (see Yoshida et al., 2012). How specific are the observed responses for 1-undecene? Can the authors use an analogous volatile chemical that does not activate AWB in the imaging and gene induction experiments?

2. The effects on foraging behavior are interesting but are a bit confusing. Do these locomotory changes lead to increased lawn-leaving behavior? The increased survival on a partial lawn of PA14 suggests they should.

3. How does the 1-undecene dependent induction of innate immune response genes relate to the survival of the worms on PA14? The authors propose that 1-undecene acts via the zip-2 pathway to induce irg-1/2/3 expression. zip-2 mutants have decreased survival on PA14 lawns, but the authors only observed a survival effect on partial lawns. Does this mean the immune response induction does not alter survival and is secondary to the behavioral response? Is survival reduced on undA mutant bacteria and does undA play a role in virulence? A major question in the field is how chemosensory and innate immune responses are coordinated to alter survival and in my opinion, this could be further fleshed out.

4. Gene expression experiments: for the RT-PCR experiments, it is unclear how many replicates were performed or how the experiments were designed. For example, it appears that Figure 1A and 1C contain the same data for irg-1/2/3, if this is the case it should be explicitly noted. If Figure 1A was essentially exploratory in nature, then in my opinion, the control data should have been replicated in order to test a different hypothesis in 1C. An alternative might be to use reporters, as in 4B, but including the appropriate controls to demonstrate an AWB specific effect. Similar controls should be performed to compare the wild-type vs. undA mutants.

Minor suggestions:

1. The chemotaxis effect in the bacterial undA mutant are striking, and it seems odd that these data are not included in the main figures.
2. There are several grammatical errors in the manuscript which could use a little more proofreading.
3. I could not find a methods entry for the acute avoidance experiments and what, if any, controls were done for this experiment?
4. Figure 3C-F all seem to show essentially the same data, perhaps this could be made more concise?
5. Line 292 makes reference to "...aversion response in at least 80% of the population" - it's unclear to what this is referencing.

Referee #2:

In this study "1-Undecene from P. aeruginosa is an olfactory signal for flight or fight response in C. elegans", the authors identified 1-undecene as a volatile product of a type of infectious
Pseudomonas aeruginosa. They showed that 1-undecene was repulsive to C. elegans worms and Pseudomonas aeruginosa unable to produce 1-undecene could not repel C. elegans. 1-undecene stimulates GCaMP3 signals in AWB neurons, and repulsion of 1-undecene to C. elegans is lost in a lim-4 mutant. Exposure to 1-undecene induces expression of several zip-2-dependent immune response genes. Overall, the authors aim to address an interesting question. The experiments are well designed, and the results are likely to contribute to a better understanding of host-pathogen interactions. Addressing or clarifying the following several questions will help to improve the manuscript.

1. The authors used an odr-3 mutant to examine the function of odor sensation in their analysis. In addition to odor sensory neurons, odr-3 is also expressed in other chemosensory neurons including nociceptive neurons ASH, phasmid neurons PHB. The results based on testing odr-3 are not fully conclusive about the involvement of odor sensation. It is helpful that authors used a lim-4 mutant to strengthen the conclusion. However, lim-4 is found in many neurons including motor neurons. The movement defects of the lim-4 mutant may interfere with chemotaxis assays. An AWB-specific ablation line or a similar reagent is needed to support the function of AWB in this study.

2. Because one single mutant was used for odr-3 and for lim-4, the conclusion based on the current results needs to be confirmed with rescue. Or, the authors should at least test a second mutation for each of these genes.

3. Figure 2b shows that when 1-undecene is tested alone, it repels worms. However, it is not clear whether 1-undecene repels worms when it is a part of the volatile products of a bacterial lawn. The authors should test whether 1-undecene added to a lawn of E. coli repels worms and quantify the lawn leaving behavior similarly as in figure 1a and 1b. In figure 2d the authors showed trajectories of worms. However, it is difficult to see on these panels where 1-undecene is and where the lawn borders are. In addition, the results need to be quantified.

4. "We observed increased speed and enhanced roaming by worms in an arena exposed to 1-undecene odor compared to the arena without the repellent (Fig. 2D and S2B). Worms also executed omega bend in response to 1-undecene in 16 out of 17 worms (Figure 2C, Supplementary Movie S3) consistent with response to an aversive signal." The speed and the omega bend need to be quantified and compared with controls.

5. T-test is perhaps not the best statistical test for the data in fig 1e, 3b and 3d.

6. Is 1-undecene a microbe-associated molecular pattern or a pathogen-associated molecular pattern? The authors propose the later, because exposure to 1-undecene induces expression of zip-2-dependent immune genes. If this is the case, mutants of PA14 with weaker virulence would not produce 1-undecene. Did author test this possibility? Clarification of this question will better characterize the property and function of 1-undecene.

7. "Moreover, we also could not detect 1-undecene in E. coli OP50, Salmonella typhimurium, and Enterococcus faecalis headspace (data not shown)". It will be very helpful to include these data.

8. It is intriguing that exposure to 1-undecene specifically upregulates zip-2-dependent immune genes irg-1,2,3. Does the induction depend on zip-2?

9. The logic for line 47-65 should be better clarified. The authors summarized several recent studies that addressed fight (immune response) and flight (avoidance and learning) responses to pathogens, odorant sensation of pathogens, virulence-independent sensation, and anatomical changes caused by pathogen exposure. It is useful for the authors to better organize these information and clarify how some or all of these studies relate to their question "However, the nature of P. aeruginosa volatile molecules that facilitate pattern recognition and threat perception in C. elegans remain unclear."

10. "However, worms have a large sensory repertoire of 302 neurons..." This sentence is not accurate.
Referee #3:

Through an analytical chemistry approach, Prakash et al. identify 1-undecene as a volatile produced by P. aeruginosa that can alter behavior and gene expression in the nematode C. elegans. They present data suggesting that it acts via the neuron AWB. The responses of C. elegans to pathogens remains an area of active research and these results will potentially interest those working in the field.

Most of the data appears robust and generally the study is well presented. There are some recent studies that should have been cited, including Ringstad's on O2/CO2 sensing, and most pertinently, "Identification of Odor Blend Used by Caenorhabditis elegans for Pathogen Recognition", Worthy et al. (2018) in the specialist journal, Chemical Senses.

The most significant claim that Prakash et al. make, "1-undecene serves as a molecular pattern and induces upregulation of a subset of immune response genes specific to P. aeruginosa in worms", is currently not sufficiently supported by the data.

To demonstrate that 1-undecene has a specific effect on host gene expression, indicative of a pathogen-specific immune response, they assayed the expression of genes that are induced by P. aeruginosa, including irg-1 and irg-2, and genes that are not induced by P. aeruginosa. Work from the Troemel lab and others, has shown that irg-1 and irg-2 are far from being pathogen-specific markers. They have been suggested to be part of an "Intracellular Pathogen Response" genes (Reddy et al 2017), regulated by pals-22 and pals-25 (Reddy et al. 2019), and induced by a range of different pathogens (e.g Orsay virus for irg-2), as a consequence of cellular dysfunction caused by infection. This is compatible with the fact irg 1 and irg 2 expression is strongly induced by the translational elongation inhibitor cycloheximide (Dunbar 2012). 1-undecene is far for a biologically inert molecule; it is known to have antimicrobial activity for example (10.3389/fmicb.2015.01082 and references therein). Its effects on host gene expression could reflect disruption of cellular homeostasis.

If the authors want to support their suggestion of a pathogen-specific response, they need to demonstrate that the 1-undecene-associated increases in gene expression are independent of pals regulation. They should also examine the expression of different markers of cellular stress. Further, they should assay the expression of candidate genes that they recently identified as being regulated by P. aeruginosa, E. faecalis and C. neoformans, such as dct-17 and lys-3.

Other minor concerns that should be addressed:

Figure 1F: N2 appear to have 2 distinct populations, one responsive and one un-responsive. There is a similar separation in Figure S2D. What is the explanation?

Welch's t-test is insensitive to equality of the variances, but does assume normal distribution. It should not be used for non-normal data (like Figure 1F).

If the control data for Figure 4C is identical to the data in Figure 4A, this needs to be indicated.

qRT-PCR is presented, for example, as "Relative mRNA expression N2 (1-Undecene / Naive)", but there are negative values, so this cannot be right. What is actually being represented?
The authors make a distinction between "younger" and "older" lawns. If 1-undecene is not produced by *P. aeruginosa* in liquid culture and/or at 37°C, the differences observed would simply be a question of the time that the bacterial cultures have been in a situation compatible with 1-undecene synthesis. This could readily checked by assaying undA expression under the different culture conditions.
Referee #1:

In this manuscript, the authors report that a Pseudomonas specific volatile odor elicits behavioral avoidance and induces immune-specific gene expression, increasing survival in response to pathogenic bacteria. There is evidence that host immune responses may be driven by chemosensory cues in multiple systems, although the molecular mechanisms of these responses are largely unknown. Recently, it was demonstrated that another C. elegans olfactory neuron - AWC - plays a critical role in non-autonomous regulation of p38 MAPK dependent immune responses via unidentified odors (Foster et al., 2020). Thus, the identification of a parallel olfactory input, including the identity of both the odors and molecular pathways involved, would be a really novel finding of broad interest.

The authors show that 1-undecene elicits behavioral avoidance and induces immune response genes and that undecene production by Pseudomonas is necessary for these responses. This is a really interesting finding, however, at this point there are many cases in which missing controls and/or an insufficiently fleshed out pathway severely limits the interpretation of these data. Additionally, it is unclear how the innate immune responses shown here are coordinated with the observed acute behavioral responses to alter survival.

Specific comments:

1. A major concern is the lack of both genetic and chemical controls for most of the experiments. For all of the C. elegans mutants, only a single allele was analyzed and no rescue experiments were performed. This is important to not only control for genetic background, but also to confirm the specificity of the proposed AWB neuron function. For example, odr-3 is expressed in AWB and AWC, as noted by the authors, but also in ASH, where it is necessary for odor responses (see Yoshida et al., 2012). How specific are the observed responses for 1-undecene? Can the authors use an analogous volatile chemical that does not activate AWB in the imaging and gene induction experiments?

RESPONSE: We agree with the reviewer that ODR-3 is also expressed non-olfactory neurons such as nociceptive neuron ASH which can respond to odors (Yoshida et al., 2012). In the revised manuscript, we have used additional allele for odr-3(n2046), in addition to odr-3(n2150) as well as lim-4(yz12) in addition to lim-4(ky403). These experiments are described in response to the next comment. We have also used ablation for ASH neurons to show that this neuron is
not involved in response to 1-undecene (Figure 3B). Additionally, we also analysed GCaMP response of ASH neurons to 1-undecene and found no response (Rebuttal Figure 1).

To control for volatile chemical stimuli, we have studied the transcriptional response of *C. elegans* upon 2 hours exposure to diacetyl (2,3 butanedione), an odor sensed by AWA olfactory neurons. We found no significant upregulation of *irg-1, irg-2* or *irg-3* (shown in rebuttal Figure 2 A and 2B) by qRT-PCR or using *irg-1::GFP* reporter.

**Rebuttal Figure 1**

![Heat map of calcium response in ASH::GCaMP3. Each row represents an individual worm recorded for 180 s under 1:100 dilution of 1-undecene, in 11 s-130 s window.]

**Rebuttal Figure 1:**
Heat map of calcium response in ASH::GCaMP3. Each row represents an individual worm recorded for 180 s under 1:100 dilution of 1-undecene, in 11 s-130 s window.

**Rebuttal Figure 2**

![Relative mRNA expression N2 (Diacetyl / Naive)]

2A  

2B

**Rebuttal Figure 2:**
(A) Real time PCR analysis of *irg-1, irg-2* and *irg-3* genes in naive and 1-undecene odor exposed N2 worms. *n* = 3. Error bars indicate SEM.
(B) *irg-1p::GFP* induction in naive worms and worms exposed to diacetyl odor (3µl, 4 spots, 2 h).

2. The effects on foraging behavior are interesting but are a bit confusing. Do these locomotory changes lead to increased lawn-leaving behavior? The increased survival on a partial lawn of PA14 suggests they should.
RESPONSE: We believe that locomotory behavior is likely a search for area with lower concentration of 1-undecene. And yes, we agree with the reviewer that avoidance is a survival strategy for worms as our experiments with partial and full lawn and with odr-3 mutants (Figure 1 in the revised manuscript) would indicate.

3. How does the 1-undecene dependent induction of innate immune response genes relate to the survival of the worms on PA14? The authors propose that 1-undecene acts via the zip-2 pathway to induce irg-1/2/3 expression. zip-2 mutants have decreased survival on PA14 lawns, but the authors only observed a survival effect on partial lawns. Does this mean the immune response induction does not alter survival and is secondary to the behavioral response? Is survival reduced on undA mutant bacteria and does undA play a role in virulence? A major question in the field is how chemosensory and innate immune responses are coordinated to alter survival and in my opinion, this could be further fleshed out.

RESPONSE: Thanks for your suggestions.

We have pre-exposed worms to 1-undecene before survival assays on *P. aeruginosa*. As shown in Figure 4 (new panel added in Figure 4F and 4G of the revised manuscript), pre-exposure to 1-undecene enhances the survival of worms from *P. aeruginosa* compared to naive worms. The experiment is shown as rebuttal Figure 3 here.

If 1-undecene is indeed a molecular pattern necessary for induction of protective immune response in worms, we expected to see reduced survival of worms on undA compared to on wild type *P. aeruginosa* PA14. As shown in rebuttal Figure 4 *C. elegans* survival is indeed reduced on undA mutant. Since we observed induction of *irg-1*, *irg-2* and *irg-3* within two hours of exposure, we would like to classify it as an immediate-early response.

**Rebuttal Figure 3**

![Rebuttal Figure 3](image)

**Rebuttal Figure 3**: Preexposure of worms to 1-undecene enhances the survival of worms upon subsequent infection with *P. aeruginosa*.

(A) Kaplan Meier survival curve on *P. aeruginosa* for N2 (naive) worms and N2 worms pre-exposed to 1-undecene odor. Survival assay was performed at 20°C.

(B) Time required for 50% of worms to die (TD$_{50}$) on *P. aeruginosa*. Each data point indicates replicates with ~100 worms. n = 3 assays. ** P ≤ 0.01 as determined by two-tailed unpaired t-test. Error bars indicate SEM.
4. Gene expression experiments: for the RT-PCR experiments, it is unclear how many replicates were performed or how the experiments were designed. For example, it appears that Figure 1A and 1C contain the same data for irg-1/2/3, if this is the case it should be explicitly noted. If Figure 1A was essentially exploratory in nature, then in my opinion, the control data should have been replicated in order to test a different hypothesis in 1C. An alternative might be to use reporters, as in 4B, but including the appropriate controls to demonstrate an AWB specific effect. Similar controls should be performed to compare the wild-type vs. undA mutants.

RESPONSE: We think the reviewer is referring to qRT-PCR analyses in Figure 4. We have taken care to describe the number of biological replicates for each experiment in the methods section of the revised manuscript. All the control data (N2 worms exposed to 1-undecene odor) has been replicated several times for the revision and included in revised Figure 4. We have ensured that control data (WT worms exposed to 1-undecene) is from different experiments across panels in Figure 4. We have also analysed the transcriptional response of 1-undecene exposure in two alleles of odr-3 and 2 alleles of lim-4 in the revised Figure 4.

Minor suggestions:

1. The chemotaxis effect in the bacterial undA mutant are striking, and it seems odd that these data are not included in the main figures.

RESPONSE: We have included the choice assay for undA mutant (over wild type PA14) in main Figure 2 in the revised manuscript.

2. There are several grammatical errors in the manuscript which could use a little more proofreading.

RESPONSE: We have proofread the entire manuscript.

3. I could not find a methods entry for the acute avoidance experiments and what, if any, controls were done for this experiment?
**RESPONSE:** For chemotaxis assays, standard protocol was used with some modifications (Bargmann et al., 1993). The details are included in the Methods section and Supplementary Figure S1D of the revised manuscript. As a control, we performed chemotaxis assays against a well-studied repellent 2-nonanone and attractant diacetyl. The data for controls is shown here (rebuttal Figure 5).

**Rebuttal Figure 5**

Chemotactic response of N2 worms towards diacetyl and 2-nonanone. Each data point indicates an individual assay plate with ~40 worms. n ≥ 3 assays. Error bars indicate SEM.

**4. Figure 3C-F all seem to show essentially the same data, perhaps this could be made more concise?**

**RESPONSE:** Thanks for your comments. Since our study is the first report on 1-undecene as a repellent and the first report of stimulation for AWB neurons, all the panels provide useful information for the *C. elegans* community. Therefore, we would like to retain all the panels.

**5. Line 292 makes reference to "...aversion response in at least 80% of the population" - it’s unclear to what this is referencing.**

**RESPONSE:** We have rephrased the statement for better understanding.

“A large fraction of worms in a population showed aversion response suggesting that 1-undecene is a physiologically relevant stimulus for worms.”
Referee #2:

In this study "1-Undecene from P. aeruginosa is an olfactory signal for flight or fight response in C. elegans", the authors identified 1-undecene as a volatile product of a type of infectious Pseudomonas aeruginosa. They showed that 1-undecene was repulsive to C. elegans worms and Pseudomonas aeruginosa unable to produce 1-undecene could not repel C. elegans. 1-undecene stimulates GCaMP3 signals in AWB neurons, and repulsion of 1-undecene to C. elegans is lost in a lim-4 mutant. Exposure to 1-undecene induces expression of several zip-2-dependent immune response genes. Overall, the authors aim to address an interesting question. The experiments are well designed, and the results are likely to contribute to a better understanding of host-pathogen interactions. Addressing or clarifying the following several questions will help to improve the manuscript.

1. The authors used an odr-3 mutant to examine the function of odor sensation in their analysis. In addition to odor sensory neurons, odr-3 is also expressed in other chemosensory neurons including nociceptive neurons ASH, phasmid neurons PHB. The results based on testing odr-3 are not fully conclusive about the involvement of odor sensation. It is helpful that authors used a lim-4 mutant to strengthen the conclusion. However, lim-4 is found in many neurons including motor neurons. The movement defects of the lim-4 mutant may interfere with chemotaxis assays. An AWB-specific ablation line or a similar reagent is needed to support the function of AWB in this study.

RESPONSE: We agree with the reviewer that ODR-3 is expressed in non-olfactory neurons such as nociceptive neuron ASH which can respond to odors. We have used ablation for ASH neurons to show that this neuron is not involved in response to 1-undecene (Figure 3B in the revised manuscript). We also analyzed GCaMP response of ASH neurons to 1-undecene and found no response (Rebuttal Figure 1).

In the revised manuscript, we have used an additional allele odr-3(n2046), in addition to odr-3(n2150). Our efforts to create AWB ablation strain have failed so far and we have not found viable ablation lines. Therefore, we have utilized an additional allele of lim-4(yz12) in addition to lim-4(ky403) and were able to phenocopy all the phenotypes.

2. Because one single mutant was used for odr-3 and for lim-4, the conclusion based on the current results needs to be confirmed with rescue. Or, the authors should at least test a second mutation for each of these genes.

RESPONSE: Thank you. We have used additional allele odr-3(n2046), in addition to odr-3(n2150) as well as lim-4(yz12) in addition to lim-4(ky403). We have included data for both the alleles of odr-3 in Figure 1 in the revised manuscript. This includes lawn leaving data (Figure 1A), survival assays (Figure 1 C-E). Transcriptional response for both alleles of odr-3 is included in Figure 4D in the revised manuscript.

We have included two alleles of lim-4 in 1-undecene chemotaxis assays (Figure 3B) and on the transcriptional response to 1-undecene exposure (Figure 4E). Both alleles fail to show induction of irg-1, irg-2 and irg-3 in response to 1-undecene.

3. Figure 2b shows that when 1-undecene is t d alone, it repels worms. However, it is not clear whether 1-uncedene repels worms when it is a part of the volatile products of a bacterial lawn. The authors should test whether 1-undecene added to a lawn of E. coli repels worms and
quantify the lawn leaving behavior similarly as in figure 1a and 1b. In figure 2d the authors showed trajectories of worms. However, it is difficult to see on these panels where 1-undecene is and where the lawn borders are. In addition, the results need to be quantified.

RESPONSE: Thanks for this suggestion. When 1-undecene was adsorbed in OP50 lawn, we could observe multiple reversals and omega turns on the lawn, as expected but we did not observe complete lawn leaving response.

We designed a different experiment to address if bacteria prefer OP50 lawn over OP50 lawn with 1-undecene. As described in the schematic in Rebuttal Figure 6, two OP50 lawns, A and B were separated by 5 mm. In test plates, 1-undecene was absorbed on lawn B or left as is in control plates. Worms were placed on lawn A and we examined if worms cross over to lawn B in 2 hours. In control plates, we found that worms moved to lawn B in all the control plates but only a small fraction of worms moved to lawn B in test plates. Based on this experiment, we inferred that C. elegans prefers OP50 food without 1-undecene.

**Rebuttal Figure 6**

(A) Schematic of the experimental setup used to record the repulsion behavior of worms from 1-undecene. Lawn A and B are E. coli OP50. Lawn B in the test condition is spiked with 0.5 µl of 1-undecene (test lawn). Worms were introduced in lawn A at the start of the experiment and the distribution of worms on lawn B was recorded after an interval of 120 m in both control and test condition.

(B) Percent distribution of worms in the test lawn (lawn B) in both control and test condition after an interval of 120 minutes. Each data point represents one assay plate with ~20 worms. n ≥ 3 assays. *** P ≤ 0.001 as determined by two-tailed unpaired t-test. Error bars indicate SEM.

4. "We observed increased speed and enhanced roaming by worms in an arena exposed to 1-undecene odor compared to the arena without the repellent (Fig. 2D and S2B). Worms also executed omega bend in response to 1-undecene in 16 out of 17 worms (Figure 2C, Supplementary Movie S3) consistent with response to an aversive signal." The speed and the omega bend need to be quantified and compared with controls.

RESPONSE: We have quantified omega turns in C. elegans exposed to 1-undecene or no odor for 5 minutes. All the worms exposed to 1-undecene odor executed omega turns (rebuttal figure 7A).

We have also quantified the speed of the worms under 1-undecene odor exposure and control condition (rebuttal figure 7B). The coordinate information of worms from 2604 tracking dataset was imported into MATLAB 2019B. The imported data was then used to calculate the distance...
by using Euclidean distance formula. Distance obtained was converted into speed by dividing with time interval between two consecutive frames. The probability density histogram was generated (bin size of 0.05) for worms’ track in the control and 1-undecene exposed conditions. Further, the significance was determined using Kolmogorov–Smirnov test. This data suggests that the average speed of worms exposed to 1-undecene was significantly higher compared to control.

**Rebuttal Figure 7**

(A) Percent of worms performing omega turn as observed under a window of 5 minutes. Each data point represents an average of 5 worms tested. n = 3. *** P ≤ 0.001 as determined by two-tailed unpaired t-test. Error bars indicate SEM.

(B) Histogram of probability density distribution of the velocity threshold of worms exposed to 1-undecene odor or control condition. Significance was calculated using Kolmogorov–Smirnov test (P = 0.0144).

5. T-test is perhaps not the best statistical test for the data in fig 1e, 3b and 3d.

**RESPONSE:** Figure 1E has been analyzed with two-tailed, unpaired t-test (Ha et al., 2010; Styer et al., 2008).

Figure 3B and 3D in the revised manuscript have been analyzed using one-way ANOVA followed by Dunnett’s multiple comparison test (Jang et al., 2019; Meisel et al., 2014; Singh and Aballay, 2019; Worthy et al., 2018).

6. Is 1-undecene a microbe-associated molecular pattern or a pathogen-associated molecular pattern? The authors propose the later, because exposure to 1-undecene induces expression of zip-2-dependent immune genes. If this is the case, mutants of PA14 with weaker virulence would not produce 1-undecene. Did author test this possibility? Clarification of this question will better characterize the property and function of 1-undecene.

**RESPONSE:** We hypothesize that *P. aeruginosa* strains with apparent increased virulence may not produce (or produce less of) 1-undecene resulting in dampened host immune response. We see reduced survival of *C. elegans* on undA mutant (rebuttal Figure 4) suggesting that 1-undecene is necessary for induction of immune response, consistent with our hypothesis. Additionally, we examined undA transcript level in a quorum-sensing defective and hypo virulent mutant *rhlR*. As shown in rebuttal Figure 8, we found no significant alteration in expression suggesting that UndA is not under the control of quorum. This suggests that there
could be several mechanisms for reduced virulence in *P. aeruginosa*, 1-undecene is just one of them and it is not altered in *rhlR* mutant.

**Rebuttal Figure 8**

Real time PCR analysis of *undA* transcript in lawn of *rhlR* mutant (24 h) over lawn of PA14 (24 h). n = 3. Error bars indicate SEM.

7. "Moreover, we also could not detect 1-undecene in *E. coli* OP50, *Salmonella typhimurium*, and *Enterococcus faecalis* headspace (data not shown)." It will be very helpful to include these data.

**RESPONSE**: Thank you. We show data for *E. coli* OP50 and *Enterococcus faecalis* OG1RF (Rebuttal Figure 9). As shown, the 1-undecene peat at 14 minutes is not seen in either of these two bacteria. We also closely surveyed the literature for several bacteria (*Staphylococcus aureus, Salmonella Typhimurium, Serratia marcescens*) and found that none of these produced 1-undecene (Siripatrawan, 2008; Worthy et al., 2018; Zhu et al., 2010).

**Rebuttal Figure 9**

9A

9B
8. It is intriguing that exposure to 1-undecene specifically upregulates zip-2-dependent immune genes irg-1,2,3. Does the induction depend on zip-2?

**RESPONSE:** Yes, the transcription of irg-1 and irg-2 is dependent in ZIP-2 transcription factor. We found that 1-undecene odor exposure for 2 hours failed to induce irg-1 and irg-2 in zip-2(tm4248) mutant. This is included in the revised manuscript as Figure 4B.

9. The logic for line 47-65 should be better clarified. The authors summarized several recent studies that addressed fight (immune response) and flight (avoidance and learning) responses to pathogens, odorant sensation of pathogens, virulence-independent sensation, and anatomical changes caused by pathogen exposure. It is useful for the authors to better organize these information and clarify how some or all of these studies relate to their question "However, the nature of P. aeruginosa volatile molecules that facilitate pattern recognition and threat perception in C. elegans remain unclear."

**RESPONSE:** We have reorganized this paragraph in the revised manuscript, to lead to the question of whether volatile molecules serve as microbe-associated molecular patterns.

10. "However, worms have a large sensory repertoire of 302 neurons..." This sentence is not accurate.

**RESPONSE:** We have rephrased to:

"However, worms have a large sensory repertoire of ~1300 G protein-coupled receptors."
Referee #3:

Through an analytical chemistry approach, Prakash et al. identify 1-undecene as a volatile produced by P. aeruginosa that can alter behavior and gene expression in the nematode C. elegans. They present data suggesting that it acts via the neuron AWB. The responses of C. elegans to pathogens remains an area of active research and these results will potentially interest those working in the field.

Most of the data appears robust and generally the study is well presented. There are some recent studies that should have been cited, including Ringstad's on O2/CO2 sensing, and most pertinently, "Identification of Odor Blend Used by Caenorhabditis elegans for Pathogen Recognition", Worthy et al. (2018) in the specialist journal, Chemical Senses.

RESPONSE: Thank you. Several additional references have been included in the revised manuscript including those mentioned by the reviewer. We have been unable to accommodate Ringstad et al, 2013 in our study as it is focused on CO2 sensing neuron and not linked to pathogen recognition or secondary metabolite perception, to the best of our knowledge.

The most significant claim that Prakash et al. make, "1-undecene serves as a molecular pattern and induces upregulation of a subset of immune response genes specific to P. aeruginosa in worms", is currently not sufficiently supported by the data.

RESPONSE: In the revised manuscript, we have pre-exposed worms to 1-undecene volatile followed by analysis of survival on live P. aeruginosa lawn. We find that pre-exposure enhances resistance of worms proving evidence that immune response induction is linked to protection (new panel added in Figure 4, 4F and 4G in the revised manuscript). The experiment is shown as rebuttal Figure 3 here. irg-1, irg-2 and irg-3 are specific to P. aeruginosa as shown previously by Troemel lab (Estes et al., 2010).

We have also analysed 8 immune effectors specific to Gram-positive bacteria and pathogenic yeast and find that they are not upregulated by exposure to 1-undecene odor (Fig. S4D). 1-undecene also does activate heat shock response, oxidative stress response (Fig. S4 G-H in the revised manuscript), or Intracellular pathogen response (rebuttal Figure 10). Collectively these experiments provide evidence that 1-undecene odor induces specific protection against P. aeruginosa and does not induce other responses.

To demonstrate that 1-undecene has a specific effect on host gene expression, indicative of a pathogen-specific immune response, they assayed the expression of genes that are induced by P. aeruginosa, including irg-1 and irg-2, and genes that are not induced by P. aeruginosa. Work from the Troemel lab and others, has shown that irg-1 and irg-2 are far from being pathogen-specific markers. They have been suggested to be part of an "Intracellular Pathogen Response" genes (Reddy et al 2017), regulated by pals-22 and pals-25 (Reddy et al. 2019), and induced by a range of different pathogens (e.g Orsay virus for irg-2), as a consequence of cellular dysfunction caused by infection. This is compatible with the fact irg 1 and irg 2 expression is strongly induced by the translational elongation inhibitor cycloheximide (Dunbar et al., 2012). 1-undecene is far for a biologically inert molecule; it is known to have antimicrobial activity for example (10.3389/fmicb.2015.01082 and references therein). Its effects on host gene expression could reflect disruption of cellular homeostasis.

RESPONSE: Thank you. Troemel lab has shown that irg-1, irg-2 and irg-3 are specific to P. aeruginosa infection and translational inhibition by the pathogen (Dunbar et al., 2012; Estes et
al., 2010). We have carefully examined the literature on IPR in C. elegans. Intracellular pathogen response is induced in worms invaded by natural viruses and Microsporidia (Reddy et al., 2017; Reddy et al., 2019). We found no evidence in the literature that irg-1, irg-2, irg-3 or zip-2 transcription are components of IPR in C. elegans. They are indeed shown to be induced only in response to P. aeruginosa (Estes et al., 2010; Troemel et al., 2006).

Additionally, we examined if 1-undecene exposure can induce IPR response by looking at an IPR reporter pals-5p::GFP, obtained from Troemel lab at UCSD. As shown in rebuttal Figure 10, 1-undecene exposure did not induce pals-5p::GFP expression. We further confirmed that P. aeruginosa live bacteria also do not induce IPR reporter pals-5 (rebuttal Figure 10). Since pals-5 was not regulated by either P. aeruginosa or 1-undecene, we did not examine upstream regulators of pals-5 such as pals-22 or pals-25. To confirm that the strain was behaving normally, we used heat shock at 30°C and found that pals-5p::GFP expression was induced in our strain as reported by Troemel lab (Reddy et al., 2017).

**Rebuttal Figure 10**

Rebuttal Figure 10: pals-5p::GFP induction in worms exposed to E. coli OP50 (blank), P. aeruginosa, E. coli OP50 under 1-undecene odor exposure and heat shock at 30°C for 24 h.

Thanks for suggestion to carefully look at the effects of the bacterial volatiles, 1-undecene, DMDS etc, on plants. It has been shown in the literature that 1-undecene in combination with another volatile promotes plant growth, although molecular mechanisms remain to be deciphered. It is especially relevant because P. aeruginosa and other Pseudomonas species are associated with plants in nature. However, in fungus, Phytophthora infestans, 1-undecene has a negative impact. In the discussion section of the revised manuscript, we have discussed the effect of 1-undecene on plants and fungi. The evidence from published literature (Hunziker et al., 2015; Lo Cantore et al., 2015) and our study point to 1-undecene as a molecular pattern which mediates interkingdom interactions. These references are included in the Discussion section of the revised manuscript.

If the authors want to support their suggestion of a pathogen-specific response, they need to demonstrate that the 1-undecene-associated increases in gene expression are independent of
pals regulation. They should also examine the expression of different markers of cellular stress. Further, they should assay the expression of candidate genes that they recently identified as being regulated by P. aeruginosa, E. faecalis and C. neoformans, such as dct-17 and lys-3.

**RESPONSE:** Please see the response to the previous comment. We found that IPR reporter *pals-5* was not upregulated by 1-undecene odor exposure (Rebuttal Figure 10). We also studied the activation of cytosolic stress response and oxidative stress response machinery upon exposure of worms to 1-undecene. As shown in rebuttal Figure 11, *hsp-16.2p*-GFP was induced by heat shock and recovery but not by 1-undecene odor exposure (Fig. S4G). Glutathione-s-transferase *gst-4p*:GFP was induced by exposure of worms to 20 mM paraquat but not by 1-undecene exposure (Fig. S4H in the revised manuscript and rebuttal Figure 12). Based on these experiments, we inferred that 1-undecene volatile does not cause disruption of cellular homeostasis.

We also examined the expression of additional genes upregulated during response of worms to *E. faecalis* and *C. neoformans*. We analysed *fmo-2, acs-2, lipl-1, lipl-3, cpr-4, cpr-5, asp-14, lys-3* and found that their transcripts were not significantly upregulated by 1-undecene odor exposure (Supplementary Figure S4D in the revised manuscript).

**Rebuttal Figure 11**

*Rebuttal Figure 11:*  
*hsp-16.2p*:GFP induction in worms exposed to *E. coli* OP50 alone (naive), OP50 with 1-undecene odor and OP50 with heat shock.

**Rebuttal Figure 12**

*Rebuttal Figure 12:*  
*gst-4p*:GFP induction in worms exposed to naive, 1-undecene odor and paraquat exposure.
**Rebuttal Figure 12:**

*gst-4p::GFP induction in worms exposed to E. coli OP50 alone (naive), OP50 with 1-undecene odor and OP50 with 20 mM paraquat.*

Other minor concerns that should be addressed:

Figure 1F: N2 appear to have 2 distinct populations, one responsive and one un-responsive. There is a similar separation in Figure S2D. What is the explanation?

**RESPONSE:** We and others have found such responses in behavioral assays. This may be linked to the expression of receptors for specific stimuli or epigenetic modification. We currently have no data to provide a solid explanation for this.

Welch’s t-test is insensitive to equality of the variances, but does assume normal distribution. It should not be used for non-normal data (like Figure 1F).

**RESPONSE:** Thanks for your suggestion. We have carefully looked at our data and most of the data follow normal distribution. Therefore, we have applied two-tailed, unpaired t-test for analyses of choice index data presented in Figure 1F in the revised manuscript. This has been used by many other groups for the analysis of choice indices (Harris et al., 2014; Pereira et al., 2020; Worthy et al., 2018).

If the control data for Figure 4C is identical to the data in Figure 4A, this needs to be indicated.

**RESPONSE:** Thank you. We have provided separate controls across panels 4A, B, D and E for qRT-PCR in the revised Figure 4.

qRT-PCR is presented, for example, as "Relative mRNA expression N2 (1-Undecene / Naive)", but there are negative values, so this cannot be right. What is actually being represented?

**RESPONSE:** The negative values are arrived at by representing FC value less than 1 as (-1/FC). This allows for better visualization of downregulation. For example, a fold change of 0.2 is a 5-fold downregulation but hard to visualize on positive axis for fold change. Thus, we use this method of representation (Dasgupta et al., 2020; Dixit et al., 2020).

The authors make a distinction between "younger" and "older" lawns. If 1-undecene is not produced by P. aeruginosa in liquid culture and/or at 37°C, the differences observed would simply be a question of the time that the bacterial cultures have been in a situation compatible with 1-undecene synthesis. This could readily checked by assaying undA expression under the different culture conditions.

**RESPONSE:** 1-Undecene is also produced in liquid culture (Rui et al., 2014; Timm et al., 2018). We have examined undA expression only on solid lawn where we study *C. elegans*-bacteria interaction and show behavioral changes. One could study the change in expression in liquid culture and change of media but that would not be relevant to our study. We also do not see an alteration in undA expression level in rhlR mutant (rebuttal Figure 8) suggesting that it may be independent of quorum sensing and population density.
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---------------------------------------END OF REBUTTAL----------------------------------------------------------
Dear Varsha,

Thank you for submitting your revised manuscript to The EMBO Journal. Your study has now been re-reviewed by the three referees and their comments are provided below.

As you can see, the referees appreciate the introduced revisions are overall supportive of the manuscript. The referees have a few more points that should be fairly straightforward to address. Regarding the point raised by referee #3 that the data doesn't sufficiently support that 1-undecene triggers a P. aeruginosa-specific response: the suggested experiment should be fairly straightforward to do or perhaps you have already done this? Let's discuss this further via video call or email.

When you submit your revised version will you also take care of the following points:

Please upload high resolution individual figures and remove the figures from the MS file. Place the figure legends at the end of the manuscript.

You can only have 5 keywords - you have at the moment 6.

Please check that there are figure callouts to Fig 4B+E and Fig S4G+H panels.

The supplementary file should be re-labelled as appendix. The figures and tables in the appendix needs to be referred to as 'Appendix Figure S#' and 'Appendix Table S#'. Please also correct callout in the text to the appendix figures/tables.

The movie legends should be removed from the appendix and ZIPed together with each movie file. The names and callouts in text needs to be corrected to 'Movie EV#'.

Methods needs correcting to Materials and Methods.

Appendix Fig S1 A panel label is missing.

Appendix Fig 3 has only one panel and so OK to refer simply as Figure 3.

We include a synopsis of the paper (see http://emboj.embopress.org/). Please provide me with a general summary statement and 3-5 bullet points that capture the key findings of the paper.

We also need a summary figure for the synopsis. The size should be 550 wide by [200-400] high (pixels). You can also use something from the figures if that is easier.

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With best wishes

Karin

Karin Dumstrei, PhD
Senior Editor
The EMBO Journal

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Referee #1:
In this revised manuscript, the authors more convincingly show that 1-undecene serves as a volatile Pseudomonas-specific pathogen associated molecular pattern. This is a significant finding of general interest. In my opinion, the authors have thoroughly addressed my concerns and I now consider this manuscript suitable for publication.

There do remain some minor proofreading or textual issues, some of which are listed below:

Calcium imaging figures do not indicate n or the number of imaging sessions.

Figure 3E - does not indicate under what imaging condition this representative image was taken.

Figure 4F - should indicate whether worms were exposed on partial or full lawns.

The statement (line 219), "We did not observe calcium response to 1-undecene in AWA, AWCon or AWCon neurons (Fig. S3A) - There appear to be small responses in AWC, so would it not perhaps be more accurate to state that you did not observe large responses in AWC?"

Referee #2:

The authors are responsive to my comments and addressed the concerns in the revised manuscript.

Referee #3:

The manuscript is much improved. I do still, however, have reservations concerning one of the authors' central claims, exemplified by the sentence in the Discussion, "The most relevant evidence for 1-undecene as a pathogen-associated molecular pattern comes from the fact that exposure of worms to volatile alone induces upregulation of immune response genes specific to P. aeruginosa".

In their rebuttal, the authors write, "irg-1, irg-2 and irg-3 are specific to P. aeruginosa as shown previously by Troemel lab (Estes et al., 2010)" and "Troemel lab has shown that irg-1, irg-2 and irg-3 are specific to P. aeruginosa infection and translational inhibition by the pathogen (Dunbar et al., 2012; Estes et al., 2010)".

Yet, as I pointed out, irg-2 expression is induced by Orsay virus, and irg-1 and irg-2 expression is strongly induced by cycloheximide. Therefore, their induction is not "specific to P. aeruginosa infection and translational inhibition by the pathogen", but to translational inhibition more generally. I would thus have expected the authors to have looked, for example, at atf-4 expression after 1-undecene exposure (by qRT-PCR).

These reservations are also due to the selective manner in which the authors interpret their data. They report a 2-fold increase for irg-1 expression as an induction (Fig. 4A), yet while the expression of both fmo-2 and acs-2 increased by more than 2-fold wrote, "We found that 1-undecene odor exposure did not induce expression of transcripts of fmo-2, acs-2, lipl-1, lipl-3, cpr-4, cpr-5, asp-14 and lys-3 (Fig. S4D)". They can't have it both ways.

Further they write, "We also analyzed the induction of heat shock response or oxidative stress response using hsp-16.2p::GFP and gst-4p::GFP respectively (Link et al., 1999; Link and Johnson,
We found that 1-undecene odor exposure did not disrupt cellular homeostasis. These results (for which the reader should be referred to FigS4G,H) are not entirely convincing, in part as the positive controls give such weak signals. Their own results for fmo-2 show how qRT-PCR is a more sensitive assay for changes in gene expression than a GFP reporter. Is there a reason why a more sensitive, qRT-PCR analysis was not performed?

As a very minimum, the authors need to tone down substantially their claim that 1-undecene is triggering a P. aeruginosa-specific response, but ideally conduct the experiments to define whether or not 1-undecene has more general effects on host physiology, by measuring for example, atf-4 mRNA levels before and after exposure.

Minor points

"qRT-PCR is presented, for example, as "Relative mRNA expression N2 (1-Undecene / Naive)", but there are negative values, so this cannot be right. What is actually being represented? RESPONSE: The negative values are arrived at by representing FC value less than 1 as (-1/FC). This allows for better visualization of downregulation. For example, a fold change of 0.2 is a 5-fold downregulation but hard to visualize on positive axis for fold change".

That's fine, but this information must be included in the figure legends!

When I wrote, "There are some recent studies that should have been cited, including Ringstad's on O2/CO2 sensing", I was referring to "Toll-like Receptor Signaling Promotes Development and Function of Sensory Neurons Required for a C. elegans Pathogen-Avoidance Behavior". Brandt JP, Ringstad N. Curr Biol. 2015. This needs to be included at, "except TOL-1 which has a limited role in C. elegans microbe interactions (Pradel et al., 2007; Tenor and Aballay, 2008)".
Referee #1:

In this revised manuscript, the authors more convincingly show that 1-undecene serves as a volatile Pseudomonas-specific pathogen associated molecular pattern. This is a significant finding of general interest. In my opinion, the authors have thoroughly addressed my concerns and I now consider this manuscript suitable for publication.

RESPONSE: Thank you.

There do remain some minor proofreading or textual issues, some of which are listed below:

Calcium imaging figures do not indicate n or the number of imaging sessions.

RESPONSE: Number of animals imaged in a single session are included in the legend for Figure 3. Two to three imaging sessions were performed for each strain.

Figure 3E - does not indicate under what imaging condition this representative image was taken.

RESPONSE: 3E represents GCaMP3 fluorescence in one of the AWB neurons 129 s after 1-undecene exposure and after 1.2 seconds (131.2 seconds on x-axis) of 1-undecene withdrawal. This information is available in the legend of Figure 3.

Figure 4F - should indicate whether worms were exposed on partial or full lawns.

RESPONSE: The worms were exposed on full lawns of P. aeruginosa.

The statement (line 219), "We did not observe calcium response to 1-undecene in AWA, AWCon or AWColf neurons (Fig. S3A) - There appear to be small responses in AWC, so would it not perhaps be more accurate to state that you did not observe large responses in AWC?"

RESPONSE: We have modified our statement to ‘We observed none or very small calcium response to 1-undecene withdrawal in AWA, AWCon or AWColf neurons’.

Referee #2:

The authors are responsive to my comments and addressed the concerns in the revised manuscript.

RESPONSE: Thank you.
Referee #3:

The manuscript is much improved. I do still, however, have reservations concerning one of the authors' central claims, exemplified by the sentence in the Discussion, "The most relevant evidence for 1-undecene as a pathogen-associated molecular pattern comes from the fact that exposure of worms to volatile alone induces upregulation of immune response genes specific to P. aeruginosa".

In their rebuttal, the authors write, "irg-1, irg-2 and irg-3 are specific to P. aeruginosa as shown previously by Troemel lab (Estes et al., 2010)" and "Troemel lab has shown that irg-1, irg-2 and irg-3 are specific to P. aeruginosa infection and translational inhibition by the pathogen (Dunbar et al., 2012; Estes et al., 2010)".

Yet, as I pointed out, irg-2 expression is induced by Orsay virus, and irg-1 and irg-2 expression is strongly induced by cycloheximide. Therefore, their induction is not "specific to P. aeruginosa infection and translational inhibition by the pathogen", but to translational inhibition more generally. I would thus have expected the authors to have looked, for example, at atf-4 expression after 1-undecene exposure (by qRT-PCR).

RESPONSE: Thanks for your suggestions. We have performed atf-4 (Glover-Cutter et al., 2013) qRT-PCR in worms exposed to 1-undecene odor. As shown in the rebuttal figure 1, we see just about 1.3-fold increase in atf-4 transcript in 1-undecene exposed worms compared to the naïve worms. We also examined an endoplasmic reticulum chaperone, hsp-4, (Shen et al., 2001) and found no significant change due to 1-undecene exposure. We have included this data in Appendix Figure S4I in the revised manuscript.

Rebuttal Figure 1

Rebuttal Figure 1:
Real time PCR analysis of transcripts for atf-4 and hsp-4 in N2 worms exposed to 1-undecene odor for 2 hours over naive N2 worms. n = 3. Error bars indicate SEM.
These reservations are also due to the selective manner in which the authors interpret their data. They report a 2-fold increase for irg-1 expression as an induction (Fig. 4A), yet while the expression of both fmo-2 and acs-2 increased by more than 2-fold wrote, "We found that 1-undecene odor exposure did not induce expression of transcripts of fmo-2, acs-2, lipl-1, lipl-3, cpr-4, cpr-5, asp-14 and lys-3 (Fig. S4D)." They can't have it both ways.

RESPONSE: Thanks for your comments. We have modified our statement in the revised manuscript to 'We found that 1-undecene odor exposure showed little or no induction in expression of transcripts of fmo-2, acs-2, lipl-1, lipl-3, cpr-4, cpr-5, asp-14 and lys-3'. Both fmo-2 and acs-2 are induced tens to hundreds of folds during infection of worms with E. faecalis or pathogenic yeast Cryptococcus neoformans (Dasgupta et al., 2020) while we see 2-3-fold increase in 1-undecene exposed worms. The 2-fold change in exposed over naive worms is rather weak in comparison to induction seen with pathogenic Gram-positive bacteria or yeast.

Further they write, "We also analyzed the induction of heat shock response or oxidative stress response using hsp-16.2p::GFP and gst-4p::GFP respectively (Link et al., 1999; Link and Johnson, 2002). We found that 1-undecene odor exposure did not disrupt cellular homeostasis". These results (for which the reader should be referred to FigS4G, H) are not entirely convincing, in part as the positive controls give such weak signals. Their own results for fmo-2 show how qRT-PCR is a more sensitive assay for changes in gene expression than a GFP reporter. Is there a reason why a more sensitive, qRT-PCR analysis was not performed?

RESPONSE: Thank you for your suggestions. We have analyzed transcript levels of hsp16.2 and gst-4 by qRT-PCR. As shown in the rebuttal figure 2, we see no significant change in the transcript levels for hsp16.2 and two-fold increase for gst-4.

Rebuttal Figure 2

Rebuttal Figure 2:
Real time PCR analysis of transcripts for hsp16.2 and gst-4 in N2 worms exposed to 1-undecene for 2 hours over naive N2 worms. n = 3. * P ≤ 0.05 as determined by two-tailed unpaired t-test with Welch’s correction. Error bars indicate SEM.
As a very minimum, the authors need to tone down substantially their claim that 1-undecene is triggering a P. aeruginosa-specific response, but ideally conduct the experiments to define whether or not 1-undecene has more general effects on host physiology, by measuring for example, atf-4 mRNA levels before and after exposure.

**RESPONSE:** qRT-PCR analyses of 8 immune effectors specific to pathogenic yeast and Gram-positive bacteria (Appendix Fig. S4D) indicated that they are either not induced at all or induced to about two folds for fmo-2 and acs-2. This is miniscule in comparison to 100-fold or higher induction seen for fmo-2 in response to infection with pathogenic bacteria (Dasgupta 2020). In addition, we find less than two-fold change in levels of transcripts for atf-4, hsp-4 and hsp16.2, and small increase in gst-4 transcript (Rebuttal Figs. 1 and 2). Microscopy reveals no increase in reporter expression for either hsp16.2 or gst-4 (Fig. S4G and S4H).

Minor points

"qRT-PCR is presented, for example, as "Relative mRNA expression N2 (1-Undecene / Naive)", but there are negative values, so this cannot be right. What is actually being represented?"

**RESPONSE:** The negative values are arrived at by representing FC value less than 1 as (-1/FC). This allows for better visualization of downregulation. For example, a fold change of 0.2 is a 5-fold downregulation but hard to visualize on positive axis for fold change”. That's fine, but this information must be included in the figure legends!

**RESPONSE:** We have included this in the legend of figure 4 in the revised manuscript.

When I wrote, "There are some recent studies that should have been cited, including Ringstad’s on O2/CO2 sensing", I was referring to "Toll-like Receptor Signaling Promotes Development and Function of Sensory Neurons Required for a C. elegans Pathogen-Avoidance Behavior". Brandt JP, Ringstad N. Curr Biol. 2015. This needs to be included at, "except TOL-1 which has a limited role in C. elegans microbe interactions (Pradel et al., 2007; Tenor and Aballay, 2008)".

**RESPONSE:** We have included the in the revised manuscript.

**REFERENCES:**

Dasgupta, M., Shashikanth, M., Gupta, A., Sandhu, A., De, A., Javed, S., and Singh, V. (2020). NHR-49 transcription factor regulates immunometabolic response and survival of Caenorhabditis elegans during Enterococcus faecalis infection. Infection and immunity 88.

Glover-Cutter, K.M., Lin, S., and Blackwell, T.K. (2013). Integration of the unfolded protein and oxidative stress responses through SKN-1/Nrf. PLoS Genet 9, e1003701.
Shen, X., Ellis, R.E., Lee, K., Liu, C.-Y., Yang, K., Solomon, A., Yoshida, H., Morimoto, R., Kurnit, D.M., and Mori, K. (2001). Complementary signaling pathways regulate the unfolded protein response and are required for C. elegans development. Cell 107, 893-903.

EDITORIAL COMMENTS

Please upload high resolution individual figures and remove the figures from the MS file. Place the figure legends at the end of the manuscript.

RESPONSE: Done.

You can only have 5 keywords - you have at the moment 5.

RESPONSE: We have included 5 keywords.

Please check that there are figure callouts to Fig 4B+E and Fig S4G+H panels.

RESPONSE: Call outs to Fig 4B+E and Fig S4G+H panels have been included in the revised manuscript.

The supplementary file should be re-labelled as appendix. The figures and tables in the appendix need to be referred to as 'Appendix Figure S#' and 'Appendix Table S#'. Please also correct callout in the text to the appendix figures/tables.

RESPONSE: We have labelled supplementary files as appendix. We have corrected callouts throughout the revised manuscript.

The movie legends should be removed from the appendix and ZIPed together with each movie file. The names and callouts in text needs to be corrected to 'Movie EV#'.

RESPONSE: Movie legends have been ZIPed together with each movie. The names and callouts have been corrected in the text.

Methods needs correcting to Materials and Methods.

RESPONSE: Corrected in the revised manuscript.

Appendix Fig S1 A panel label is missing.

RESPONSE: Corrected.

Appendix Fig 3 has only one panel and so OK to refer simply as Figure 3.

RESPONSE: Corrected.
We include a synopsis of the paper (see http://emboj.embopress.org/). Please provide me with a general summary statement and 3–5 bullet points that capture the key findings of the paper.

RESPONSE:

Summary Statement: Well-studied pathogen-associated molecular patterns (PAMPs) are constituents of microbial cells such as certain components of their cell wall. Here, we show that a volatile molecule produced by Pseudomonas aeruginosa activates a pathogen-specific immune response in Caenorhabditis elegans host. This study presents a new paradigm for pathogen recognition resulting in activation of flight or fight response.

• 1-undecene volatile produced by P. aeruginosa activates flight response in C. elegans on short time scale of second to minutes.
• Longer exposure of worms to 1-undecene induces a pathogen-specific immune response.
• Both flight and fight response induced by 1-undecene rely on the AWB olfactory neurons of worms.
• Preexposure of worms to 1-undecene protects them against a subsequent infection with P. aeruginosa.
• 1-Undecene serves as a pathogen-associated molecular pattern (PAMP) for C. elegans.

We also need a summary figure for the synopsis. The size should be 550 wide by [200-400] high (pixels). You can also use something from the figures if that is easier.

RESPONSE: We have uploaded a summary figure.
Dear Varsha,

Thank you for submitting your revised manuscript. I have now had a chance to take a look at it and I appreciate the introduced changes.

I am therefore very pleased to accept the manuscript for publication here.

Congratulations on a nice study

Best Karin

Karin Dumstrei, PhD
Senior Editor
The EMBO Journal

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### A. Figures

#### 1. Data

The data shown in figures should satisfy the following conditions:

- The data were obtained and processed according to the field’s best practice and are presented to reflect the results of the experiments in an accurate and unbiased manner.
- Figure panels include only data points, measurements or observations that can be compared to each other in a scientifically meaningful way.
- Graphs include clearly labeled error bars for independent experiments and sample sizes. Unless justified, error bars should not be shown for technical replicates.
- If n = 5, the individual data points from each experiment should be plotted and any statistical test employed should be justified.
- Source Data should be included to report the data underlying graphs. Please follow the guidelines set out in the author shop guidelines on Data Presentation.

#### 2. Captions

Each figure caption should contain the following information, for each panel where they are relevant:

- a specification of the experimental system investigated (e.g. cell line, species name).
- the assay(s) and method(s) used to carry out the reported observations and measurements
- an explicit mention of the biological and chemical entity(ies) that are altered/varied/perturbed in a controlled manner.
- the exact sample size (n) for each experimental group/condition, given as a number, not a range;
- an explicit mention of the biological and chemical entity(ies) that are being measured.
- a statement of how many times the experiment shown was independently replicated in the laboratory.
- definitions of statistical methods and measures:
  - common tests, such as t-test (please specify whether paired vs. unpaired), simple χ² tests, Wilcoxon and Mann-Whitney tests, can be unambiguously identified by name only, but more complex techniques should be described in the methods section;
  - are tests one-sided or two-sided?
  - are tests one-sided or two-sided?
  - are there adjustments for multiple comparisons?
  - exact statistical test results, e.g., F-values = n but not p-values = n;
  - definition of ‘center values’ as median or average;
  - definition of error bars as s.e.m. or s.d.

Any descriptions too long for the figure legend should be included in the methods section and/or with the source data.

### B. Statistics and general methods

#### 1. How was the sample size chosen to ensure adequate power to detect a pre-specified effect size?

| Sample size was determined based on similar assays described by other investigators in the field. |

#### 2. For animal studies, include a statement about sample size estimate even if no statistical methods were used.

| Sample size was determined based on similar assays described by other investigators in the field. |

#### 3. Describe inclusion/exclusion criteria if samples or animals were excluded from the analysis. Were the criteria pre-established?

| No animals were excluded from the analysis of behaviors described in this study. |

#### 4. Were any steps taken to minimize the effects of subjective bias when allocating animals/samples to treatment (e.g., randomization procedure)? If yes, please describe.

| No |

#### 5. Are there any steps to minimize the effects of subjective bias during group allocation or/and when assessing results (e.g. binding of the investigator)? If yes please describe.

| No randomization was used. |

#### 6. For animal studies, include a statement about randomization even if no randomization was used.

| No randomization was used. |

#### 7. For animal studies, include a statement about blinding even if no blinding was done.

| No blinding was performed for C. elegans. |

#### 8. For every figure, are statistical tests justified as appropriate?

| Yes |

#### 9. Do the data meet the assumptions of the tests (e.g., normal distribution)? Describe any methods used to assess it.

| GraphPad PRIZM was used to assess distribution of the data. |

#### 10. Is there an estimate of variation within each group of data?

| Yes |
G- Dual use research of concern

18. Provide a “Data Availability” section at the end of the Materials & Methods, listing the accession codes for data generated in this study and deposited in a public database (e.g. PRIDE https://www.ebi.ac.uk/pride/archive/) or a public registry (e.g. BioProject number and/or clone number, supplementary information or reference to an antibody validation profile. e.g., Antibodypedia (see link list at top right), IDPRG waive (see link list at top right).

C- Reagents

1. Identify the source of all reagents and report if they were recently authenticated (e.g., by iTRAQ profiling) and tested for mycoplasma contamination. Not applicable

D- Animal Models

8. Report species, strain, gender, age of animals and genetic modification status where applicable. Please detail housing and husbandry conditions and the source of animals. Yes

9. For experiments involving live vertebrates, include a statement of compliance with ethical regulations and identify the committee(s) approving the experiments. Not applicable

10. We recommend consulting the ARRIVE guidelines (see link list at top right) (Powell B, S.I., et al., 2010) to ensure that other relevant aspects of animal studies are adequately reported. See author guidelines, under “Reporting Guidelines”. See also: NIH (see link list at top right) and MRC (see link list at top right) recommendations. Please confirm compliance. Yes

E- Human Subjects

15. Report the clinical trial registration number (at ClinicalTrials.gov or equivalent), where applicable. Not applicable

16. For phase II and III randomized controlled trials, please refer to the CONSORT flow diagram (see link list at top right) and submit the CONSORT checklist (see link list at top right) with your submission. See author guidelines, under “Reporting Guidelines”. Please confirm you have submitted this list. Not applicable

17. For human subject research, we recommend that you follow the REMARK reporting guidelines (see link list at top right). See author guidelines, under “Reporting Guidelines”. Please confirm you have followed these guidelines. Not applicable

F- Data Accessibility

18. Provide a “Data Availability” section at the end of the Materials & Methods, listing the accession codes for data generated in this study and deposited in a public database (e.g. RNA-Seq data: Gene Expression Omnibus GSE139492, Proteomics data: PRIDE PXD000208 etc.) Please refer to our author guidelines for ‘Data Deposition’. Statement included

20. Access to human clinical and genomic datasets should be provided with as few restrictions as possible while respecting ethical obligations to the patients and relevant medical and legal issues. If practicable possible and compatible with the individual consent agreement used in the study, such data should be deposited in one of the major public access-controlled repositories such as dbGAP (see link list at top right) or EGA (see link list at top right). Not applicable

21. Computational models that are central and integral to a study should be shared without restrictions and provided in a machine-readable form. The relevant accession numbers or links should be provided. When possible, standardized formats (BML, CellML) should be used instead of scripts (e.g., MATLAB). Authors are strongly encouraged to follow the MIBML guidelines (see link list at top right) and submit their model in a public database such as BioModels (see link list at top right) or EGA (see link list at top right). If computer source code is provided with the paper, it should be deposited in a public repository or included in supplementary information. Not applicable

G- Dual use research of concern

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