Abl Kinase Inhibits the Engulfment of Apoptotic Cells in Caenorhabditis elegans


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The engulfment of apoptotic cells is required for normal metazoan development and tissue remodeling. In *Caenorhabditis elegans*, two parallel and partially redundant conserved pathways act in cell-corpse engulfment. One pathway includes the adaptor protein CED-2 CrkII and the small GTPase CED-10 Rac, and acts to rearrange the cytoskeleton of the engulfing cell. The other pathway includes the receptor tyrosine kinase CED-1 and might recruit membranes to extend the surface of the engulfing cell. Although many components required for engulfment have been identified, little is known about inhibition of engulfment. The tyrosine kinase Abl regulates the actin cytoskeleton in mammals and *Drosophila* in multiple ways. For example, Abl inhibits cell migration via phosphorylation of CrkII. We tested whether ABL-1, the *C. elegans* ortholog of Abi, inhibits the CED-2 CrkII-dependent engulfment of apoptotic cells. Our genetic studies indicate that ABL-1 inhibits apoptotic cell engulfment, but not through CED-2 CrkII, and instead acts in parallel to the two known engulfment pathways. The CED-10 Rac pathway is also required for proper migration of the distal tip cells (DTCs) during the development of the *C. elegans* gonad. The loss of ABL-1 function partially restores normal DTC migration in the CED-10 Rac pathway mutants. We found that ABL-1 is the *C. elegans* homolog of mammalian Abi (Abi interactor) proteins, is required for engulfment of apoptotic cells and proper DTC migration. Like Abi, Abi proteins are cytoskeletal regulators. ABL-1 acts in parallel to the two known engulfment pathways, likely downstream of ABL-1. ABL-1 and ABI-1 interact physically in vitro. We propose that ABL-1 opposes the engulfment of apoptotic cells by inhibiting ABI-1 via a pathway that is distinct from the two known engulfment pathways.

**Introduction**

Regulated reorganization of the cytoskeleton is a fundamental process in tissue morphogenesis and physiologic cell migration [1]. Dysregulation of the cytoskeleton is a hallmark of pathologic processes, such as cancer cell invasion and metastasis [2]. The engulfment of apoptotic cells requires a major cytoskeletal reorganization within the engulfing cell, which must extend its plasma membrane completely around the dying cell. In *C. elegans*, neighboring cells engulf apoptotic cells. Eleven genes appear to act in two parallel pathways required for engulfment: ced-1, ced-6, ced-7, and dyn-1; and ced-2, ced-5, ced-10, ced-12, mig-2, unc-73, and pdr-1 (Figure 1) [3]. These two pathways have been proposed to recruit membranes for cell surface extension and rearrange the cytoskeleton, respectively. The pathways together promote the extension of the engulfing cell around the apoptotic cell.

In the pathway for membrane recruitment, which we refer to as the CED-1 pathway (see below), four proteins have been identified (Figure 1). CED-7 is an ABC transporter required in both the engulfing cell and the engulfed cell and might expose a pro-engulfment signal on the surface of the apoptotic cell [4,5]. The role of CED-7 in the engulfing cell has not been defined. CED-7 is thought to signal through CED-1, a receptor on the engulfing cell surface homologous to *Drosophila* Draper and the mammalian EGF-like receptor MEGF10 [6]. CED-1, in turn, is proposed to signal through CED-6, a protein that contains a phosphotyrosine-binding domain [7]; CED-6 can bind a motif in the intracellular domain of CED-1 [8] and is thought to activate DYN-1, a *C. elegans* dynamin homolog [9]. DYN-1, by analogy to its role in vesicular transport in mammalian cells, might recruit membrane for the engulfment process. The CED-1 pathway also is involved in degrading apoptotic cells once they are engulfed [10,11].

The pathway for cytoskeletal rearrangement requires the small GTPase CED-10 Rac, and we refer to this pathway as the CED-10 Rac pathway. Two parallel pathways contribute to CED-10 Rac activation (Figure 1). CED-2, the *C. elegans* homolog of the oncprotein CrkII, is an SH2 and SH3 domain-containing adaptor protein [12] that interacts with an atypical heterodimeric guanine nucleotide exchange factor (GEF) consisting of the proteins CED-5 [13] and CED-12 [14–16], homologs of mammalian DOCK180 and ELMO, respectively. In mammals, a signal from the apoptotic

**Abbreviations:** Abi, Abi interactor; DIC, differential interference contrast; DTC, distal tip cell; GEF, guanine nucleotide exchange factor; SH, Src homology

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Author Summary

Cell death or apoptosis is a normal part of animal development, as is the engulfment and removal of dead cells by other cells. In the nematode Caenorhabditis elegans, ten highly conserved proteins have been characterized previously for their roles in engulfment and in cell migration, both of which involve the formation of cellular extensions. Little is known, however, about how engulfment is inhibited. In mammals, the tyrosine kinase Abl, which regulates the actin cytoskeleton and which when misexpressed causes two types of leukemia, prevents the CrkII protein from facilitating cell migration. CrkII functions in engulfment in C. elegans and mammals. We tested whether the C. elegans homolog of Abl, ABL-1, could inhibit engulfment. We found that ABL-1 functions as an inhibitor of apoptotic cell engulfment and cell migration. However, our analysis further showed that ABL-1 does not function by inhibiting other known engulfment proteins, including C. elegans CrkII. Our data indicate that ABL-1 blocks ABI-1, the C. elegans homolog of the mammalian and Drosophila Abl-interactor (Abi) cytoskeletal- regulatory proteins. We propose that ABL-1 acts via ABI-1 to inhibit a newly identified pathway during cell corpse engulfment and cell migration.

cell to the engulfing cell is transduced through CrkII to the DOCK180/ELMO heterodimer [17], and an analogous process is thought to occur between CED-2 and the CED-5/CED-12 heterodimer. The CED-5/CED-12 GEF activates the Rac1 homolog CED-10, and activated CED-10 rearranges the cytoskeleton [18,19]. Rac proteins are members of the Rho family of small GTPases that regulate the cytoskeleton and function in intracellular signaling [20]. The phosphatidylinositol receptor PSR-1, which recognizes phosphatidylinositol on the surface of the dying cell, has been proposed to act upstream of CED-2 [21].

MIG-2, the mammalian homolog of RoG, another Rho family GTPase also regulates the CED-10 Rac pathway [22]. MIG-2 acts on CED-5/CED-12 in parallel to CED-2 [23]. UNC-73, a RoGEF homologous to the mammalian protein Trio, activates MIG-2 [23]. The MIG-2 branch of the CED-10 Rac pathway provides a minor input into the engulfment pathway: mutations in mig-2 and unc-73 enhance the defects of other engulfment mutants but do not cause engulfment defects on their own [23].

Although many of the proteins that act in these two engulfment pathways are known, how these pathways are regulated is poorly understood. In mammals, the Abl tyrosine kinase functions in multiple processes that regulate the actin cytoskeleton [24]. For example, Abl blocks cell migration by phosphorylating and inhibiting the CED-2 homolog CrkII [25]. In addition to its role in Rac-dependent cell migration, Abl has been implicated in multiple signaling pathways in both the cytoplasm and the nucleus [26]. Abl acts in numerous cell biological processes, including cytoskeletal rearrangement, cell migration, apoptosis, transcription, and the response to oxidative stress [24,26]. Dysregulation of Abl via fusion of the Abl gene to the BCR gene is the cause of chronic myelogenous leukemia (CML) and an aggressive subtype of acute lymphoblastic leukemia (ALL) [27,28]. Recently, Abl signaling has been implicated in the prevention of breast cancer tumorigenesis [29] and in pathological fibrosis caused by the chemotherapeutic agent bleomycin [30,31].

That Abl functions in multiple diverse biological processes is reflected by its complex domain structure. In addition to its tyrosine kinase and DNA-binding domains, Abl has several protein-binding domains: Src Homology 2 (SH2), SH3, and F- and G-actin binding domains. Over 70 Abl-interacting proteins have been identified, largely through biochemical studies and cell-culture experiments. The in vivo relevance of most of these interactions has not been conclusively determined.

The C. elegans genome encodes a single Abl homolog, ABL-1. As in other organisms, Abl in C. elegans has many functions. ABL-1 protects germline cells from programmed cell death in response to ionizing radiation by antagonizing a molecular pathway that contains cell cycle checkpoint proteins and the p53 homolog CEP-1 [32]. ABL-1 is required for Shigella flexneri pathogenesis through an unknown mechanism [33]. A function for ABL-1 in cytoskeletal regulation has also been described: ABL-1 regulates epidermal morphogenesis in the C. elegans embryo by opposing the Ena/VASP homolog UNC-34 [34].

We found that ABL-1 inhibits the engulfment of apoptotic cells. Our genetic studies indicate that ABL-1 acts independently of both known engulfment pathways, suggesting the
existence of another pathway for engulfment. We show that ABI-1, the *C. elegans* homolog of the Abi (Abl interactor) cytoskeletal and signaling family of proteins, is a member of this newly identified pathway.

**Results**

**ABL-1 Inhibits the Engulfment of Apoptotic Cell Corpses**

To test whether abl-1 has a role in engulfment, we counted the number of unengulfed apoptotic cell corpses in the heads of first larval stage (L1) animals harboring mutations in abl-1 and engulfment pathway genes. The number of unengulfed corpses varies with the strength of the engulfment defect and defines a quantitative assay of engulfment defects [35]. We used two presumptive null alleles of *abl-1* in this study, *n*1963 and *ok*171. *n*1963 is a G-to-A transition at the splice acceptor of exon 10 (bp 16967 of the M79 cosmid sequence), resulting in removal of most of the kinase domain and the change of a conserved arginine to serine. *ok*171 is a deletion allele that defines a quantitative assay of engulfment defects [35]. We tested for interactions between *abl-1* and the genes of the CED-10 Rac pathway. *abl-1* mutation did not modify the engulfment defects of either *ced-5(n*1812) or *ced-12(n*3261) null mutants (Table 1). *abl-1* function also did not modify the engulfment defect of *ced-2(n*1994); *mig-2(gm*38 *mu*133) null double mutants, in which both known inputs into the CED-10 Rac pathway are absent (Figure 1). However, *abl-1* mutation partially suppressed the engulfment defect caused by *ced-2(n*1994) alone (the number of cell corpses decreased from 22.0 to 15.7, *p* < 0.0001) and by the partial loss-of-function allele *ced-2(e*1752) alone (a decrease from 18.9 to 13.2, *p* < 0.0001). *mig-2* null mutations do not cause engulfment defects on their own so they were tested in combination with a CED-1 pathway mutant (see below). Animals completely lacking *ced-10* die as embryos and were not tested. ABL-1 function did not depend on the presence of functional CED-1, CED-6, or CED-7 and therefore ABL-1 acts independently or downstream of the CED-1 pathway. These data are consistent with a role for ABL-1 in the negative regulation of apoptotic cell engulfment. Experiments that address alternative explanations for the affect of ABL-1 on engulfment are presented in the next section.

We tested for interactions between *abl-1* and the genes of the CED-10 Rac pathway. *abl-1* mutation did not modify the engulfment defects of either *ced-5(n*1812) or *ced-12(n*3261) null mutants (Table 1). *abl-1* mutation also did not modify the engulfment defect of *ced-2(n*1994); *mig-2(gm*38 *mu*133) null double mutants, in which both known inputs into the CED-10 Rac pathway are absent (Figure 1). However, *abl-1* mutation partially suppressed the engulfment defect caused by *ced-2(n*1994) alone (the number of cell corpses decreased from 22.0 to 15.7, *p* < 0.0001) and by the partial loss-of-function allele *ced-2(e*1752) alone (a decrease from 18.9 to 13.2, *p* < 0.0001). *mig-2* null mutations do not cause engulfment defects on their own so they were tested in combination with a CED-1 pathway mutant (see below). Animals completely lacking *ced-10* die as embryos and were not tested, but the engulfment defect caused by a partial loss-of-function allele, *ced-10(n*1993), was suppressed by *abl-1(0)* (the number of cell corpses decreased from 20 to 8.7, *p* < 0.0001).

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**Table 1. abl-1 Mutations Suppress the Engulfment Defects of Engulfment ced Gene Mutations**

<table>
<thead>
<tr>
<th>Genotype</th>
<th>n Corpses ± sd</th>
<th>n</th>
<th>p-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild-type</td>
<td>0 ± 0</td>
<td>10</td>
<td>—</td>
</tr>
<tr>
<td><em>abl-1(n</em>1963)*</td>
<td>0 ± 0</td>
<td>10</td>
<td>—</td>
</tr>
<tr>
<td><em>ced-1(n</em>2091)*</td>
<td>22.1 ± 4.2</td>
<td>20</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td><em>ced-1(n</em>2091); <em>abl-1(n</em>1963)*</td>
<td>13.7 ± 3.5</td>
<td>20</td>
<td>—</td>
</tr>
<tr>
<td><em>ced-1(e</em>1735)*</td>
<td>25.5 ± 4.5</td>
<td>21</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td><em>ced-1(n</em>1735); <em>abl-1(n</em>1963)*</td>
<td>19.5 ± 4.3</td>
<td>21</td>
<td>—</td>
</tr>
<tr>
<td><em>ced-2(e</em>1752)*</td>
<td>18.9 ± 3.0</td>
<td>21</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td><em>ced-2(e</em>1752); <em>abl-1(n</em>1963)*</td>
<td>13.2 ± 3.4</td>
<td>20</td>
<td>—</td>
</tr>
<tr>
<td><em>ced-2(n</em>1994)*</td>
<td>22.0 ± 4.3</td>
<td>21</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td><em>ced-1(e</em>1735); <em>ced-2(n</em>1994)*</td>
<td>15.7 ± 3.1</td>
<td>20</td>
<td>—</td>
</tr>
<tr>
<td><em>ced-1(e</em>1735); <em>ced-5(n</em>1812)*</td>
<td>16.7 ± 3.4</td>
<td>20</td>
<td>—</td>
</tr>
<tr>
<td><em>ced-5(n</em>1812)*</td>
<td>32.9 ± 4.0</td>
<td>16</td>
<td>&gt;0.2</td>
</tr>
<tr>
<td><em>ced-5(n</em>1812); <em>abl-1(n</em>1963)*</td>
<td>32.2 ± 6.4</td>
<td>19</td>
<td>—</td>
</tr>
<tr>
<td><em>ced-6(n</em>2095)*</td>
<td>26.8 ± 5.4</td>
<td>24</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td><em>ced-6(n</em>2095); <em>abl-1(ok</em>171)*</td>
<td>19.6 ± 5.1</td>
<td>20</td>
<td>—</td>
</tr>
<tr>
<td><em>ced-7(n</em>1892)*</td>
<td>35.6 ± 4.5</td>
<td>19</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td><em>ced-7(n</em>1892); <em>abl-1(n</em>1963)*</td>
<td>29.9 ± 2.2</td>
<td>20</td>
<td>—</td>
</tr>
<tr>
<td><em>ced-10(n</em>1993)*</td>
<td>20.0 ± 3.6</td>
<td>23</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td><em>ced-10(n</em>1993); <em>abl-1(n</em>1963)*</td>
<td>8.7 ± 3.2</td>
<td>20</td>
<td>—</td>
</tr>
<tr>
<td><em>ced-10(n</em>1993); <em>ok</em>171</td>
<td>8.3 ± 3.3</td>
<td>20</td>
<td>—</td>
</tr>
<tr>
<td><em>ced-12(tp</em>2)*</td>
<td>15.3 ± 4.6</td>
<td>21</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td><em>ced-12(tp</em>2); <em>abl-1(n</em>1963)*</td>
<td>6.4 ± 3.5</td>
<td>20</td>
<td>—</td>
</tr>
<tr>
<td><em>ced-12(n</em>3261)*</td>
<td>22.7 ± 5.3</td>
<td>20</td>
<td>&lt;0.1</td>
</tr>
<tr>
<td><em>ced-12(n</em>3261); <em>abl-1(n</em>1963)*</td>
<td>20.4 ± 4.3</td>
<td>20</td>
<td>—</td>
</tr>
<tr>
<td><em>ced-1(e</em>1735); <em>ced-2(n</em>1994)*</td>
<td>35.5 ± 4.6</td>
<td>19</td>
<td>&gt;0.2</td>
</tr>
<tr>
<td><em>ced-1(e</em>1735); <em>ced-2(n</em>1994); <em>abl-1(n</em>1963)*</td>
<td>34.5 ± 3.7</td>
<td>19</td>
<td>—</td>
</tr>
<tr>
<td><em>ced-1(n</em>2091); <em>mig-2(gm</em>38 <em>mu</em>133)*</td>
<td>24.0 ± 4.0</td>
<td>22</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td><em>ced-1(n</em>2091); <em>abl-1(n</em>1963) <em>mig-2(gm</em>38 <em>mu</em>133)*</td>
<td>18.3 ± 3.6</td>
<td>22</td>
<td>—</td>
</tr>
<tr>
<td><em>ced-2(n</em>1994); <em>mig-2(gm</em>38 <em>mu</em>133)*</td>
<td>32.7 ± 4.1</td>
<td>21</td>
<td>&gt;0.2</td>
</tr>
<tr>
<td><em>ced-2(n</em>1994); <em>abl-1(n</em>1963) <em>mig-2(gm</em>38 <em>mu</em>133)*</td>
<td>32.4 ± 8.8</td>
<td>20</td>
<td>—</td>
</tr>
</tbody>
</table>

First larval stage (L1) worms were anaesthetized and viewed using DIC microscopy. The numbers of cell corpses in the heads were counted. sd, standard deviation.

*This p-value refers to comparisons between the strain with the engulfment mutation alone and the strains with the engulfment mutation with either the *abl-1(n*1963) or the *abl-1(ok*171) mutation.

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We also tested whether the engulfment defect of a ced-1(e1735); ced-2(n1994) double mutant could be suppressed. A mutation in abl-1 did not modify the engulfment defect of the double mutant, even though each single mutant was suppressed. This result is consistent with the possibility that ABL-1 acts upstream of both the CED-10 Rac and the CED-1 pathways. Alternatively, ABL-1 might suppress a pathway parallel to these two pathways, but its suppression might be too weak to modify an engulfment defect as severe as that of the ced-1(e1735); ced-2(n1994) double mutant. We present data below that support the latter model.

Because mig-2 and unc-73 mutations enhance the engulfment defects of other engulfment gene mutations but do not cause defects on their own [23], we tested whether ABL-1 acts through the Mig-2 branch of the CED-10 Rac pathway by testing whether an abl-1 mutation could suppress the mig-2 enhancement of ced-1(n2091). We observed fewer apoptotic cell corpses in the heads of ced-1(n2091); mig-2(gm38 mu133) animals when an abl-1 mutation was present (Table 1), demonstrating that ABL-1 can act in the absence of Mig-2 function. Therefore, ABL-1 does not act solely through either the CED-2 branch or the Mig-2 branch of the CED-10 Rac engulfment pathway.

In summary, abl-1(1f) suppressed partial but not complete loss of the CED-10 Rac pathway. The inability of abl-1(1f) to suppress the engulfment defects of the CED-10 Rac pathway when this pathway was completely nonfunctional (i.e., ced-5 null, ced-12 null, or ced-2; mig-2 double null mutants, Figure 1) suggests that the CED-10 Rac pathway genes do not function by blocking the action of ABL-1. Instead, ABL-1 might inhibit the CED-10 Rac pathway, or ABL-1 might signal in parallel to the CED-10 Rac pathway through another group of effectors that require CED-10 Rac pathway function to accomplish apoptotic cell engulfment.

Notably, the observation that abl-1(1f) suppressed the engulfment defect of mutants that completely lack CED-2 function indicates that the effect of ABL-1 on engulfment is at least partially independent of CED-2, i.e., ABL-1 does not act only by inhibiting CED-2. Therefore, *C. elegans* ABL-1 can act to inhibit CED-2 CrkII-dependent pathways via a mechanism distinct from the known mechanism in mammals, in which Abl phosphorylates the CED-2 homolog CrkII.

### ABL-1 Does Not Affect the Cell-Death Process Directly

The effect of *abl-1* mutation on the number of unengulfed corpses could be caused by mechanisms other than the direct inhibition of cell engulfment. *abl-1* mutation might (1) suppress programmed cell death, resulting in fewer cell corpses; (2) alter the timing of corpse appearance during development, resulting in fewer corpses at the time of observation; or (3) change cell-corpuses so that they were not recognized as corpses or were unstable and were lost altogether.

To determine whether *abl-1* acts in programmed cell death, we evaluated whether cells known to undergo programmed cell death did so normally. In wild-type animals, 16 cells undergo programmed cell death in the anterior pharynx during embryogenesis [36], and their deaths can be scored by direct observation of their nuclei using DIC microscopy [35]. Mutants defective in programmed cell death, such as mutants with null mutations in the caspase *ced-3*, have up to 14 extra recognizable cell nuclei in the anterior pharynx [35,37]. We observed no extra nuclei in either *abl-1*(n1693) or *abl-1*(ok171) animals (Table 2). We also used a more stringent test for cell-death defects: enhancement of the death defect of *ced-3(n2427)* mutants, which are partially defective in programmed cell death [38]. As shown in Table 2, *ced-3(n2427)* animals had an average of 1.7 extra corpses, and *abl-1* mutation did not enhance the *ced-3(n2427)* death defect.

In the ventral nerve cord, six Pn.aap cells (P1.aap, P2.aap, and P9–12.aap) undergo programmed cell death postembryonically in wild-type animals, but not in death-deficient mutants [38,39]. Defects in programmed cell death are easily detected and quantified for five of these cells (P2.aap, P9–P12.aap) using a *lin-11::gfp* transcriptional reporter transgene, which is expressed by surviving Pn.aap cells [38]. As in the pharynx, in the ventral nerve cord *abl-1* mutation neither caused excess cell survival on its own nor enhanced the death defect caused by the *ced-3(n2427)* mutation (Table 2). We conclude that ABL-1 did not promote programmed cell death of non-germline cells. ABL-1 does protect against programmed cell death in the germline, most notably after radiation exposure [32]. In the germline *abl-1* loss-of-function therefore causes excess cell death, not suppression of cell death. If this function of ABL-1 were present in non-germline cells, we would expect increased numbers of corpses, not the reduction of unengulfed corpses that we observed.

To test whether ABL-1 affects the timing, persistence, or morphology of cell corpses, we used time-lapse DIC microscopy to observe wild-type and *abl-1*(n1693) embryos for 150 min after the first appearance of a cell corpse [9]. During this time, approximately 70 cell corpses appear in the wild-type animal. We observed no significant difference between wild-type animals and *abl-1* mutants with respect to the number of corpses that appeared or when they appeared (Figure 2A). Also, the length of time that corpses persisted was similar between wild-type and *abl-1*(n1693) animals (Figure 2B). In addition, apoptotic cell corpses in *abl-1*(n1693) animals looked identical to wild-type corpses (Figure 2C). We conclude that loss of ABL-1 did not affect the time of appearance or morphology of apoptotic cell corpses.

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**Table 2. abl-1 Mutation Does Not Block Cell Death**

<table>
<thead>
<tr>
<th>Tissue</th>
<th>Genotype</th>
<th>n Extra Cells</th>
<th>n</th>
<th>sd</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pharynx</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wild-type</td>
<td></td>
<td>0.0 ± 0.2</td>
<td>33</td>
<td></td>
</tr>
<tr>
<td>ced-3(n2436)</td>
<td></td>
<td>6.1 ± 1.5</td>
<td>16</td>
<td></td>
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<tr>
<td>ced-3(n1717)</td>
<td></td>
<td>10.8 ± 1.4</td>
<td>14</td>
<td></td>
</tr>
<tr>
<td>abl-1(n1693)</td>
<td></td>
<td>0 ± 0</td>
<td>28</td>
<td></td>
</tr>
<tr>
<td>abl-1(ok171)</td>
<td></td>
<td>0 ± 0</td>
<td>5</td>
<td></td>
</tr>
<tr>
<td>ced-3(n2427)*</td>
<td></td>
<td>1.7 ± 1.1</td>
<td>10</td>
<td></td>
</tr>
<tr>
<td>ced-3(n2427); abl-1(n1693)*</td>
<td></td>
<td>1.4 ± 1.0</td>
<td>17</td>
<td></td>
</tr>
<tr>
<td>ced-3(n2427); abl-1(ok171)</td>
<td></td>
<td>1.1 ± 1.1</td>
<td>7</td>
<td></td>
</tr>
<tr>
<td>Ventricle nerve cord</td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Wild-type</td>
<td></td>
<td>0.0 ± 0.2</td>
<td>25</td>
<td></td>
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<tr>
<td>abl-1(n1693)*</td>
<td></td>
<td>0.0 ± 0.2</td>
<td>25</td>
<td></td>
</tr>
<tr>
<td>ced-3(n2427)*</td>
<td></td>
<td>2.3 ± 1.3</td>
<td>100</td>
<td></td>
</tr>
<tr>
<td>ced-3(n2427); abl-1(n1693)*</td>
<td></td>
<td>2.1 ± 1.1</td>
<td>100</td>
<td></td>
</tr>
</tbody>
</table>

Third larval stage (L3) worms were anaesthetized and viewed using DIC microscopy. The number of extra cell nuclei in the anterior pharynxes were counted. sd, standard deviation.

*These strains contained nIs96 (lin-11::gfp) V.

doi:10.1371/journal.pbio.1000099.t002
**abl-1 Mutation Suppresses Other Defects Associated with Engulfment Pathway Genes**

Mutants defective in corpse engulfment are also partially defective in programmed cell death, indicating that cell-corpses engulfment promotes cell killing [38,40]. A role for engulfment genes in promoting programmed cell death has also been found in *Drosophila* [41], showing that the pro-apoptotic function of cell engulfment is evolutionarily conserved.

In *C. elegans* mutants partially defective in cell-killing (e.g., *ced-3* caspase partial loss-of-function mutants), some cells that are fated to die undergo some of the morphological changes that accompany programmed cell death but then recover and persist as normal cells [38,40]. Most cells fated to die will nonetheless die. In *C. elegans* mutants with a partial loss of *ced-3* function and a mutation in an engulfment gene, a much larger number of cells fated to die will survive.

We tested whether *abl-1* mutations suppress the cell-death defect caused by engulfment gene mutations. The anterior pharynges of animals doubly mutant for *ced-3(n2427)*, a partial loss-of-function mutation, and a mutation in an engulfment gene, with or without *abl-1(n1963)* were scored for the presence of extra nuclei (Figure 3A). Partial loss-of-function and null mutations in *ced-1* and *ced-12* were analyzed. We observed fewer extra nuclei in animals that had the *abl-1(n1963)* mutation with either of the *ced-1* mutations and with the partial loss-of-function *ced-12(tp2)* mutation (Figure 3A). For
example, *ced-1(e1735); ced-3(n2427)* animals had an average of 5.9 extra cells in their pharynges, whereas *ced-1(e1735); ced-3(n2427); abl-1(n1963)* animals had an average of 4.2 extra cells (*p < 0.001*). However, *abl-1* mutation did not affect the number of extra cells seen in *ced-12(tp2); ced-3(n2427)* animals. These findings are consistent with our observations concerning the role of ABL-1 in engulfment: *abl-1* mutation suppressed the death defect of a null mutation in the CED-1 pathway but did not suppress the death defect of a null mutation in the CED-10 Rac pathway (*ced-12(n3261)*).

We also examined the effect of an *abl-1* mutation on engulfment gene-related death defects in the ventral nerve
ABL-1 acts in parallel to or downstream of the CED-10 Rac pathway to do so.

We also examined the effect of a loss of abl-1 function on the DTC migration defect of an animal harboring the ced-10(n3417) deletion mutation, a putative ced-10 null allele [22]. Because the ced-10(n3417) mutation causes maternal-effect lethality (i.e., homozygous null animals produce no live progeny), we analyzed the homozygous progeny of ced-10 heterozygotes (ced-10(n3417)/lin-1(e1275) dpy-13(e184sd)) with or without the abl-1(ok171) deletion mutation. These ced-10(n3417) homozygous animals presumably survived because they have CED-10 protein derived from maternally provided ced-10 mRNA. As with ced-2, ced-5, and ced-12 null mutants, the DTC defect of these ced-10 null mutants was suppressed by an abl-1 loss-of-function mutation: the percentage of defective gonadal arms in ced-10(n3417) animals decreased from 20.7% to 7.6% in ced-10(n3417); abl-1(ok171) animals (p < 0.005) (Figure 4). We note that these ced-10 animals are unlikely to totally lack ced-10 function, since the ced-10 null phenotype is maternal-effect lethal, indicating that ced-10 homozygotes derived from ced-10+ heterozygotes have some ced-10 function; that ced-10+ heterozygotes indeed have some ced-10 function is supported by the observation that only 20.7% of the DTCs of the ced-10(n3417) animals migrated inappropriately, which is far less than that seen in ced-5(n1812), ced-12(n3261), or ced-10(n1993) animals. Therefore, no compelling conclusion about whether ABL-1 acts in parallel to or downstream of the CED-10 Rac pathway can be made on the basis of this experiment.

Notably, ced-10(n3417); abl-1(ok171) animals produced a small number of live progeny (unlike ced-10(n3417) animals), some of which achieved adulthood; none of these was fertile. This suppression of the maternal-effect lethality caused by ced-10(n3417) might well reflect an effect of the abl-1(ok171) mutation in the complete absence of ced-10 function. If so, at least in this case, abl-1 acts in parallel to or downstream of ced-10. We suggest it is simplest to postulate that abl-1 also acts in parallel to or downstream of ced-10 for engulfment, engulfment-mediated programmed cell death, and DTC migration.

Our findings concerning engulfment and gonadal migration are consistent with two models of ABL-1 function. In one model, ABL-1 acts through different molecular pathways to inhibit the morphological changes that drive engulfment of apoptotic cells and to inhibit the migration of DTCs, i.e., ABL-1 acts directly on CED-10 or another protein in the CED-10 Rac pathway in engulfment and on a different set of proteins in DTC migration. Alternatively, ABL-1 acts in a pathway distinct from the CED-10 Rac pathway but common to both processes, and this common pathway is more important in DTC migration than in engulfment. For example, in gonadal cell migration, either the CED-10 Rac pathway or a second ABL-1-inhibited pathway might be sufficient for normal DTC migration. If this were the case, loss of ABL-1 function would derepress the ABL-1-regulated pathway and suppress DTC migration defects even in the absence of any CED-10 Rac pathway function, as we observed. In engulfment, the requirement for the CED-10 Rac pathway might not be able to be overcome by derepression of the ABL-1-regulated pathway. We present data below supporting the second model, namely that ABL-1 acts in a common pathway distinct from the CED-10 Rac and CED-1 pathways in engulfment and gonadal cell migration.
ABL-1 Probably Acts in Engulfing Cells

To determine whether ABL-1 acts in the engulfing or the engulfed cell, we performed ectopic expression experiments in which abl-1 was expressed from a transgene in an abl-1 mutant background. Specifically, we used a protocol adapted from Reddien and Horvitz [12] and expressed abl-1 under the control of C. elegans heat-shock promoters at a time at which all embryonic deaths are complete; we then counted the number of embryonic cell corpses in the heads of first larval stage (L1) animals. Any rescue that occurs cannot involve transgene function in the engulfed cells, which have already died. All cells in the head that die during embryogenesis do so prior to 5 h before hatching, and we scored engulfment within 5 h of heat shock. We found that expression of abl-1 in ced-10(n1993); abl-1(n1963) animals increased the number of unengulfed corpses in L1 heads from 9.8 to 18.2 (Table 3) (for comparison, ced-10(n1993) animals had 20.0 corpses, Table 1). abl-1 expression almost completely abrogated the effect of the abl-1(n1963) mutation, whereas expression of a gfp-only control transgene had minimal effects on the engulfment (Table 3). We also tested whether ABI-1, the only Abi protein in C. elegans, affected engulfment (Figure 5A). C. elegans ABI-1 is 31% identical to its closest human homolog, ABI-1 (Figure 5B), and has higher conservation in predicted functional domains. Our determination of abl-1 transcript structures is described in Text S1.

We tested ABL-1 function using the deletion mutation tm494, which removes a 66-amino acid region just before the SH3 domain and changes the frame of the remaining sequence, resulting in a C-terminal truncation just before the SH3 domain. We also used abi-1 RNAi by feeding to reduce abi-1 expression [63,64], because RNAi by injection [65] causes embryonic lethality [66]. We showed that abi-1(tm494) and feeding abi-1 RNAi probably caused a very weak loss of function, given that animals were viable and fertile. abi-1(tm494) or abi-1 RNAi in wild-type animals caused a weak engulfment defect (Table 4). Also, the tm494 mutation or abi-1 RNAi significantly enhanced the engulfment defects of all engulfment mutants tested (in both the CED-10 Rac and CED-1 pathways) (Table 4). In addition, abi-1 RNAi significantly enhanced the engulfment defect of a ced-1(e1735); ced-5(n1812) double mutant, in which both engulfment pathways are nonfunctional. To determine whether ABI-1 function requires ABL-1, we assayed the effect of abi-1 RNAi in strains doubly mutant for engulfment genes and abl-1. Loss of abi-1 function enhanced the engulfment defects of these strains to the same degree, whether abl-1 mutation was present or absent. Therefore, ABI-1 does not act by modulating ABL-1 function.

We also assayed the effect of abi-1 RNAi on gonadal migration. abi-1 RNAi enhanced the ced-5 defect in gonadal cell migration (from 47.5% to 58.8%, p < 0.04) (Figure 6). Strikingly, abi-1(lf) completely abrogated the effect of abl-1 mutation on gonadal cell migration in ced-5 and ced-12 animals.

Because Abi proteins are in the Scar/WAVE complex and other Scar/WAVE complex members have been implicated in engulfment (GEX-2 and GEX-3, the C. elegans homologs of Sra-1 and Nap, respectively) [67], we asked whether the localization of GEX-3 is altered in animals with a loss of abi-1 function. We assessed the localization of a rescuing

<table>
<thead>
<tr>
<th>Table 3. Overexpression of abl-1 Reverses the Effect of abl-1 (n1963) on Engulfment in ced-10(n1993); abl-1(n1963) Animals</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Transgene</strong></td>
</tr>
<tr>
<td>----------------</td>
</tr>
<tr>
<td>P(hsp-gfp)</td>
</tr>
<tr>
<td>P(hsp-gfp)</td>
</tr>
<tr>
<td>P(hsp-abl-1)</td>
</tr>
<tr>
<td>P(hsp-abl-1)</td>
</tr>
<tr>
<td>P(hsp-abl-1(K340R))</td>
</tr>
<tr>
<td>P(hsp-abl-1(K340R))</td>
</tr>
</tbody>
</table>

ced-10(n1993); abl-1(n1963) embryos containing the transgenes indicated above were heat shocked at 33 °C for 1 h and then allowed to recover for 3–3.5 h at 20 °C. The numbers of cell corpses in the heads of anaesthetized first larval stage (L1) worms were counted using DIC microscopy within 0.5 h of hatching. The number of persistent corpses was determined from two independent transgenic lines for each transgene. sd, standard deviation.

doi:10.1371/journal.pbio.1000099.003
GFP::GEX-3 fusion protein (kindly provided by M. Soto) in embryos of animals that contain a gex-3 null mutation (zu196) with either a wild-type abl-1 allele or abl-1(ok171). We found no localization of the fusion protein around embryonic cell corpses and no apparent difference in the pattern of fluorescence in both strains (Figure S1). Therefore, Scar/WAVE complexes do not appear to localize around apoptotic cell corpses at least at the time that corpses become visible by DIC optics. Furthermore, the absence of ABL-1 protein does not appear to alter the localization of Scar/WAVE complexes. However, ABL-1 probably only interacts with a fraction of the Scar/WAVE complexes present, and this fraction might not be sufficiently large to detect a difference in the overall localization of Scar/WAVE complexes using this method. Moreover, the rescuing GFP::GEX-3 fusion transgene is located on an extrachromosomal array and might be expressed at much higher levels than wild-type GEX-3. For these reasons, we cannot unambiguously interpret these results.

In summary, loss of ABI-1 function enhanced the engulfment defects caused by the inactivation of either the CED-10 Rac pathway or the CED-1 pathway or both together and enhanced the DTC migration defects of the CED-10 Rac pathway.

Figure 5. abi-1 Is a C. elegans Abi Gene

(A) Alignment of three Homo sapiens Abi proteins (Abi-1, Abi-2, and Abi-3) with C. elegans ABI-1. The dashed line indicates the Wave-binding domain. The dotted line shows the proline-rich region. The continuous line shows the SH3 domain. The boxed residues in ABI-1 indicate the sequence removed by the tm494 deletion. Dark gray indicates identities, light gray similarities.

(B) Similarity and identity indices between ABI-1 and each of the three human Abi proteins.

doi:10.1371/journal.pbio.1000099.g005

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In summary, loss of ABI-1 function enhanced the engulfment defects caused by the inactivation of either the CED-10 Rac pathway or the CED-1 pathway or both together and enhanced the DTC migration defects of the CED-10 Rac pathway.
ABL-1 and ABI-1 Interact In Vitro

In mammals, Abl and Abi-2 interact in vitro in two ways. The SH3 domain of Abl binds to a site in the first 157 amino acids of Abi-2, likely a proline-rich site. The SH3 domain of Abi-2 binds to a proline-rich region near the center of Abl (amino acids 593–730) [49]. To test whether C. elegans ABL-1 and ABI-1 interact directly, we performed in vitro binding experiments with glutathione-S-transferase fused to the N terminus of ABI-1 and in vitro translated portions of ABL-1. We made two ABL-1 constructs, an N-terminal fragment (amino acids 112–611) and a C-terminal fragment (amino acids 606–1,224). ABL-1(112–611) bound to ABI-1, but to a small degree also bound to GST alone. Quantification of the bands using phosphorimagery revealed 8-fold higher binding in the ABI-1 lane than in the GST lane despite a much smaller amount of ABI-1 than GST loaded on the gel (note Coomassie Blue-stained gel next to the autoradiograph) (Figure 7). ABL-1 did not bind to the control Luciferase; we also failed to observe ABL-1(606–1,224) binding. ABL-1::gfp reporter broadly throughout embryogenesis (unpublished data).

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Table 4. abi-1 Mutation Enhances the Engulfment Defects of Engulfment Genes

<table>
<thead>
<tr>
<th>Genotype</th>
<th>n Corpses ± sd</th>
<th>n</th>
<th>p-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild-type</td>
<td>2.9 ± 0.2</td>
<td>18</td>
<td>&lt;0.05*</td>
</tr>
<tr>
<td>wild-type + abi-1 RNAi</td>
<td>4.6 ± 0.5</td>
<td>19</td>
<td>—</td>
</tr>
<tr>
<td>ced-1(e1735)</td>
<td>32.9 ± 3.2</td>
<td>21</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>ced-1(e1735) + L4440 RNAi</td>
<td>32.9 ± 3.2</td>
<td>21</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>ced-1(e1735) + abi-1 RNAi</td>
<td>22.0 ± 4.1</td>
<td>21</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>ced-1(e1735) + ced-2(n1994)</td>
<td>22.0 ± 4.1</td>
<td>21</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>ced-2(n1994) + L4440 RNAi</td>
<td>32.9 ± 3.2</td>
<td>21</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>ced-2(n1994) + abi-1 RNAi</td>
<td>32.9 ± 3.2</td>
<td>21</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>ced-2(n1994) + ced-5(n1812)</td>
<td>32.9 ± 3.2</td>
<td>21</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>ced-5(n1812) + L4440 RNAi</td>
<td>32.9 ± 3.2</td>
<td>21</td>
<td>&lt;0.0001</td>
</tr>
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<td>21</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

First larval stage (L1) worms were anesthetized and viewed using DIC microscopy. The number of cell corpses in the heads were counted. sd, standard deviation.

*p-Value refers to comparison between wild type and abi-1 RNAi. Comparison between wild type and abi-1(tm494) did not reach statistical significance.

doi:10.1371/journal.pbio.1000099.g004

The engulment pathway. ABI-1 action was not modified by ABL-1 inactivation. These data suggest that ABL-1 acts through ABI-1 to inhibit engulfment and DTC migration. abi-1 is expressed broadly throughout embryogenesis [32]. To determine where abi-1 is expressed, we created an abi-1::gfp reporter transgene and observed expression of the abi-1 reporter broadly throughout embryogenesis (unpublished data.).

Engulfment Pathway Genes

The SH3 domain of ABI-1 and in vitro translated portions of ABL-1. We made two ABL-1 constructs, an N-terminal fragment (amino acids 112–611) and a C-terminal fragment (amino acids 606–1,224). ABL-1(112–611) bound to ABI-1, but to a small degree also bound to GST alone. Quantification of the bands using phosphorimagery revealed 8-fold higher binding in the ABI-1 lane than in the GST lane despite a much smaller amount of ABI-1 than GST loaded on the gel (note Coomassie Blue-stained gel next to the autoradiograph) (Figure 7). ABL-1 did not bind to the control Luciferase; we also failed to observe ABL-1(606–1,224) binding. ABL-1::gfp reporter broadly throughout embryogenesis (unpublished data).

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Discussion

We have demonstrated that the *C. elegans* Abl ortholog ABL-1 negatively regulates the engulfment of apoptotic cells. *abl-1* inhibits the engulfment process as well as the engulfment-related cell-killing process and the migration of DTCs during gonadogenesis. Our genetic analysis suggests that ABL-1 acts in a manner that does not require the known engulfment pathways. Ectopic expression experiments indicate that ABL-1 acts in engulfing cells and that its function at least partially depends on its kinase activity. Moreover, our studies show that the Abi ortholog ABI-1 acts in engulfment and DTC migration. Finally, our genetic and biochemical studies both suggest that ABL-1 directly inhibits ABI-1 in a pathway distinct from the known engulfment gene pathways.

How Do ABL-1 and ABI-1 Interact In Vivo?

Our genetic analysis indicates that either *abl-1* could inhibit *abi-1* or the two genes could act in separate molecular pathways. We favor the former hypothesis, both because ABL-1 and ABI-1 interact directly in vitro and because in mammalian cell culture and cultured neurons the homologs of these proteins interact and function in processes that regulate the cytoskeleton [55,56].

Our findings establish that an Abl protein and an Abi protein interact functionally in vivo. In some in vitro studies Abi activated Abl [49,51–54], in others Abi activated Abi [56,57], and in still others Abi appeared to block Abl function [48–50,55]. We found that *abl-1* and *abi-1* have opposing functions in vivo. Given the large number of proteins with which Abl and Abi interact and the multiple cellular contexts in which they function, in vivo analyses will be critical to distinguish which interactions are relevant for a particular cell biological process.

ABL-1/ABI-1 Likely Act in Parallel to the CED-10 Rac and CED-1 Pathways

The inhibition of engulfment by ABL-1 occurred in the absence of functional CED-2 CrkII, indicating that the effect of ABL-1 on engulfment and DTC migration is not mediated by CED-2 CrkII inhibition. In mammals, Abl phosphorylates tyrosine 221 of CrkII between its SH3 domains, resulting in inhibition of CrkII function and suppression of cell migration [25,68]. This tyrosine is not conserved in *C. elegans*. We conclude that *C. elegans* ABL-1 blocks the CED-10 Rac pathway by a novel mechanism.

Our analysis of genetic interactions between *abl-1* and the engulfment genes suggests the existence of a new pathway involved in both cell-corpse engulfment and DTC migration. Loss of ABL-1 function suppressed the engulfment and cell-death defects of all CED-1 pathway genes tested. *abl-1* mutation did not suppress the engulfment defects of *ced-5* or *ced-12* nulls but did suppress their DTC migration defects. Since *abl-1* mutation modulated DTC migration in the absence of *ced-5* or *ced-12* function (i.e., when the CED-10 Rac pathway was inactive), ABL-1 can signal through another pathway. We propose that ABL-1 acts in a third pathway not only for DTC migration but also for engulfment. If so, this pathway cannot promote engulfment in the absence of CED-10 Rac activation. Although it is formally possible that the function of ABL-1 in engulfment is mediated through the CED-10 Rac pathway while its effect on DTC migration is mediated through a different pathway, we prefer a simpler model in which ABL-1 acts through a single pathway to oppose the engulfment genes. This pathway might act in parallel to CED-10 or it might act on CED-10 Rac (see below).

We found that ABI-1 is required for the function of ABL-1 in engulfment and DTC migration. ABL-1 promoted engulfment and migration independently of the known engulfment pathways downstream of or in parallel to ABL-1. We propose that ABL-1 inhibits ABI-1 and that these two proteins define a new molecular pathway required for cell-corpse engulfment and DTC migration. Interestingly, loss of *abl-1* did not suppress the engulfment defects of CED-10 Rac pathway null mutants, whereas loss of *abi-1* did enhance those engulfment defects. At least three models can explain these findings. ABL-1 might not be a sufficiently potent inhibitor of ABI-1 to
Figure 8. ABL-1 and ABI-1 Likely Function in Parallel to the CED-10 Rac and CED-1 Engulfment Pathways

We suggest that ABL-1 inhibits ABI-1, which acts to promote the engulfment of apoptotic cells. ABL-1 might signal either independently of the CED-10 Rac pathway (Arrow 1) or through CED-10 Rac, in parallel to the CED-10 GEF CED-5/CED-12 (Arrow 2). This model also applies to the roles of ABL-1 and ABI-1 in the regulation of DTC migration, but we do not show DTC migration in the figure, because the CED-1 pathway does not act in this process. Since ABL-1 and ABI-1 act in DTC migration, ABL-1/ABI-1 cannot act solely through the CED-1 pathway since the proteins of the CED-1 pathway have no role in DTC migration.

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affect engulfment in the absence of CED-10 Rac pathway activity. Second, the CED-10 Rac pathway might be absolutely required for engulfment, so that derepressing the ABL-1/ABI-1 pathway does not suppress the CED-10 Rac pathway engulfment defect. Third, loss of abl-1 function might increase engulfment activity in the absence of the CED-10 Rac pathway insufficiently to detect in the engulfment assay. Loss of abl-1 function suppressed the DTC migration defect of a null ced-10 mutant. However, these animals were the progeny of heterozygotes and probably contained some C. elegans ced-10 of a null mutation. This observation provides stronger evidence that abl-1 can act in parallel to or downstream of ced-10. We were unable to test the effects of abl-1 or abi-1 mutations on a null mutant of dyn-1, because this mutant arrests development during embryogenesis and, unlike in the case of a null ced-10 mutant, DTC migration is not affected in dyn-1 mutants. Therefore, we do not know whether ABL-1 or ABI-1 can act independently of CED-10 or DYN-1 for engulfment or engulfment-mediated programmed cell death (or DTC migration in the case of CED-10) and hence whether ABL-1 and ABI-1 act through either of these genes or in a parallel molecular pathway. We think it unlikely that ABL-1 and ABI-1 act through DYN-1, since the CED-1 pathway has no known role in DTC migration, and ABL-1 and ABI-1 modulate DTC migration defects.

There are at least three models for how the ABL-1 and ABI-1 proteins act in engulfment and DTC migration. First, ABL-1 might directly inhibit ABI-1 from promoting engulfment of apoptotic cells and inappropriate DTC migration through a molecular pathway that acts in parallel to the known engulfment gene pathways (Figure 8). Second, ABL-1 might act on CED-10 Rac in parallel to the CED-5/CED-12 heterodimer (Figure 8). These models are not mutually exclusive. In a third model, the CED-10 Rac pathway and ABL-1 both act on ABI-1 in parallel to each other, with CED-10 Rac activating ABI-1 and ABL-1 inhibiting it. Studies of mammalian Abl proteins are consistent with the first two models. For example, Abl proteins are found in complexes with N-WASP [60] and the Formins [59]. Both N-WASP and formins act in actin cytoskeletal rearrangements independently of the CED-10 homolog Rac. C. elegans Abl-1 might act similarly in our first model. Abl proteins also form a complex with Eps8 and Sos-1. Formation of the Abl-1/Eps8/Sos-1 complex activates the RacGEF activity of Sos-1 in response to tyrosine kinase signaling [50,61]. ABI-1 might act this way in our second model. In mammals, Rac and Abl proteins both act on the Scar/WAVE complex through interactions with Abl proteins [53,69]. However, in these cases Abl activates Abl. By contrast, we found that ABL-1 inhibits ABI-1. For this reason we do not favor a model in which ABL-1 and CED-10 Rac act on ABI-1 in parallel.

Why Negatively Regulate Engulfment?

Despite the large number of genes known to be involved in engulfment, only a few negatively regulate the process. Loss of SWAN-1, a CED-10-binding protein, suppresses the engulfment and DTC migration defects caused by ced-10 loss of function, and SWAN-1 thus might be a negative regulator of these processes [70]. In mammalian macrophages and macrophage cell lines, the small GTPase Rho and one of its effectors, Rho-kinase, negatively regulate engulfment of apoptotic cells [71,72]. Rho, like Rac, regulates the cytoskeleton, and in many contexts the two proteins act in opposition to each other [20]. All Rac proteins are downregulated by Rac-specific GTPases (RacGAPs) [73]. No RacGAPs have been identified that function in engulfment, though presumably one or more will be found.

Pathways that inhibit engulfment might prevent the inappropriate engulfment of healthy cells that are not programmed to die. There are examples of inappropriate engulfment of mammalian cells. In entosis, cells engulf and eventually kill neighboring cells that have lost their attachments to the extracellular matrix [74]. Also, the glycosylated surface protein SIRPα is found on engulfing cells and interacts with the integrin-associated protein CD47 on other viable cells. When that interaction is disrupted, inappropriate engulfment occurs [75]. Notably, the intracellular cascades that transduce these signaling events have not been discovered. Possibly abl-1 transduces these types of signals.

Perhaps there are conditions that cause cells to be particularly sensitive to engulfment, and without such negative regulatory pathways cells would be inappropriately killed. Since engulfment promotes the cell-killing process, several of the situations in which engulfment occurs could be severely affected by such excess cell death: organismoal development, wound healing and infection control.

Targeting signaling pathways that negatively regulate engulfment might have therapeutic benefits. For example, inducing professional engulfing cells, such as macrophages, to engulf diseased cells, such as cancerous cells or those infected with viruses or bacteria, could aid in combating these disease processes. Also, in humans impaired engulfment of apoptotic...
cells has been associated with systemic lupus erythematosus (SLE) [76] and in mice, ineffective engulfment of apoptotic cells can cause an SLE-like syndrome [77]. Enhancing the engulfment of apoptotic cells might aid in treating or preventing certain autoimmune disorders. Extremely effective and specific small molecule inhibitors of Abl, such as Imatinib [78] (Gleevec) and Nilotinib [79], exist so this idea could be tested.

Materials and Methods

Strains and genetics. *C. elegans* strains were maintained at 20 °C as described [80]. The N2 Bristol strain was used as the wild-type strain. Animals were grown on NGM plates and fed OP50 bacteria [42]. The mutations and integrants used were: LGL, cad-1(e2735), unc-24(e1275), unc-3(e138); LGX, 1(e1275), unc-24(e138), ced-2(e1752, n1994), ced-3(n717, n2424, n2427, n2436), ced-12(n3261, tp2), des-1(e1752), unc-24(e138); LGV, unc-3(e566), gmIn144, unc-66(e7169), lin-1(e1275); unc-24(e138); LGV, unc-3(e566), gm114, unc-66(e7169); lin-1(e1275). Mutant alleles for which no citation is given were described previously [81]. Information about 2°/tm alleles kindly provided by S. Moudy (Women’s Medical University, Japan) can be found at www.wormbase.org. The following balancer chromosomes were used: LGI; LGIII, hT2[qIs48]; LGIV, mIn1[mIs14]; LGIV, LGV, nT1[qIs51].

Isolation of abl-1 alleles. abl-1 alleles (gm327, gm332, n1961, n1963, n1966) were isolated in two separate genetic screens for mutations that suppressed the Unc locomotion phenotype of unc-34 mutants. unc-34 mutant *C. elegans* were mutagenized with ethyl methanesulfonate (EMS) as described [80]. The F2 progeny of mutagenized unc-34(e566) hermaphrodites representing 17,500 haploid genomes were inspected for the presence of animals with normal locomotion. Three strains were identified as containing candidate suppressors (n1961, n1963, n1964) [82], gm327 and gm332 were isolated in an F1 clonal screen. unc-34(gm114) P0 hermaphrodites were mutagenized and allowed to lay eggs. Single F1 animals were transferred onto fresh plates. P1 progeny were screened for a non-Unc phenotype. 10,952 haploid genomes were screened this way.

In addition to the five alleles of abl-1, five other mutations were isolated in the above screens. Complementation and mapping studies demonstrated that these five mutations defined a second complementation group and are alleles of the gene called-1 [83]. See Text S1 for a more detailed description of the complementation tests and mapping and the cloning of the abl-1 gene.

Quantitation of engulfment defects. Unengulfed apoptotic corpses were visualized in the heads of young larvae as refractile discs directly using Nomarski DIC microscopy with a compound microscope (Thornwood). For animals treated with feeding RNAi, L1 animals were picked, and those with gonads that had not passed the four-cell stage were viewed as described above. P-values for pairwise comparisons were calculated using the Student’s t-test.

Quantitation of cell-death defects. For quantitation of cell-death defects in the anterior pharynx, animals in the third larval stage (L3) were anesthetized and viewed with DIC microscopy as described above. Briefly, the locations of the nuclei of the 16 cells that undergo programmed cell death in the anterior pharynx are known [36]. In wild-type animals by the L3 stage, all of those nuclei have disappeared; any remaining nuclei in the animals tested were scored as extra cells. P-values for pairwise comparisons in the pharynges were calculated using Student’s t-test. For quantitation of cell-death defects in the ventral nerve cord, a dissecting microscope equipped with an ultraviolet light source (Kramer Scientific) was used. To analyze GFP expression by the L3 stage, all of those nuclei have disappeared; any remaining nuclei in the animals tested were scored as extra cells. P-values for pairwise comparisons between strains with wild-type abl-1 and mutant abl-1, we tested the null hypothesis that the median number of extra VC neuron-like cells in the wild-type strain was less than or equal to one more than the median number of VC neuron-like cells in the mutant strain separately, as conducted with the wilcox.test function of R (www.r-project.org).

Time-lapse microscopy. Gravid *C. elegans* were dissected, and embryos at the two-cell stage were placed at 20 °C for 180 min. Single embryos were then placed on an agar pad, sealed with petroleum jelly, and viewed at 22 °C using a Zeiss Axioscope 2 compound microscope equipped with Nomarski DIC accessories, a Hamamatsu ORCA-ER digital camera, and Openlab image acquisition software (Improvision). Pictures were taken every 3 min for 200 min, starting 180 min after the first cell division. The time of appearance of each component of each embryo first, at least 4 serial z sections at 0.5 μm section were recorded. Recording began at 180 min after first cleavage, because the first apoptotic cell corpses appear at approximately 200 min after the first cell division [36]. Images were analyzed with Imaris (www.bitplane.com) and ImageJ (rsbweb.nih.gov/ij); the plugins Manual Tracking (rsbweb.nih.gov/ij/plugins/track/track.html) and Ome Locci (http://www.loci.wisc.edu/ome). P-values for comparisons between strains were calculated using the Wilcoxon rank-sum test.

Quantitation of DTC defects. Adult animals 18 h after the mid-fourth larval stage (L4) were anesthetized and viewed as described above. "Quantitation of engulfment defects," and gonads were visualized [85,86]. Only gonads that were completely visualized were scored. DTC migration was scored as defective when the gonad was morphologically abnormal (extra turn, two arms, or bizarre twists) or when the gonad was short or long. Gonadal length was defined as abnormal when the gonad tip was distal to the ipsilateral sperma-tetheca (short) or distal to the contralateral spermatheca (long). The vast majority of abnormalities in morphology rather than in length. P-values for pairwise comparisons were calculated by Fisher’s exact test.

abl-1 rescue. *P*<sub>shgabl-1</sub> and *P*<sub>shgabl-1</sub>(K340R) were constructed as follows. The entire coding sequence (bp 1–3,675) of abl-1 was synthesized by PCR from an *abl-1* cDNA template (yk14820, kindly provided by M. Kohara). The PCR product was cloned into the vector pCR8/GW/TOPO (TOPO TA cloning kit; Invitrogen) that had been generated using the Quickchange Site-Directed Mutagenesis kit (Stratagene) from the pCR8GW/TOPO plasmid containing the *abl-1* PCR product described above using the following primers: forward: 5'-GCCACATGACGCAATTTGATCTGGCATGGATTGAAAGAGCATGATGCC-3'; reverse: 5'-GCGATTGATCTTCTTC-CAACGGTGGAGATCCGAGTACCATGCGTGAG-3'. These primers created the K340R mutation. DNA sequences of PCR products were determined for accuracy and orientation. *abl-1* products were then cloned into pDEST-MB1 and pDEST-MB7 (kindly provided by M. Boxem) using the Gateway method (Invitrogen). pDEST-MB1 and pDEST-MB7 are the plasmids pPR49.78 and pPR49.83, respectively, with Gateway cassettes inserted, allowing for Gateway cloning. The *P*<sub>shgabl-1</sub> plasmids have been described previously [12]. *P*<sub>shgabl-1</sub>, *P*<sub>shgabl-1</sub>(K340R), and *P*<sub>shgabl-1</sub> plasmids were injected into *cgl-1(e1963); abk-1(e364)* animals at a concentration of 20 ng/μl with a plasmid containing myo-2::gfp as a coinjection marker at 5 ng/μl and with 35 ng/μl of 1 Kb Plus DNA ladder (Invitrogen) for a total concentration of 80 ng/μl per injection. The pharyngeal and transgenic animals were RP-positive. Embryos were grown at 20 °C, heat shocked at 25 °C for 20 min, and then cell corpses were counted in the heads of newly hatched first larval stage (L1) animals. Two independent transgenic lines were analyzed for each transgene combination.

Expression analysis of the *abl-1* promoter. A PCR product containing the *abl-1* promoter (622 bp of DNA encoding the sequence between the 5’ end and the adjacent gene start [B0336.5]) followed by the *gfp* gene coding sequence was constructed as described in Wormbook (http://www.wormbook.org/chapters/ www região refusions reportergenusions.html). The *gfp* gene encoding GFP (pSF6565) and vector pRF4 (52), both from M. Chalfie. *C. elegans* kit (Addgene). The PCR product was injected into the gonads of animals at a concentration of 10 ng/μl with a plasmid containing the unc-76 gene (p76-1B9) [87] as a coinjection marker at 5 ng/μl and with 1 Kb Plus DNA ladder (Invitrogen) at 30 ng/μl. Three independent non-Unc transgenic lines were analyzed using fluorescence microscopy and DIC microscopy.

RNA interference by feeding. Animals were fed bacteria that contained either the RNAi empty feeding vector L4440 [63] or an RNAi feeding vector with part of the *abl-1* gene, B0336.6, cloned into the vector. Bacteria were cultured as described previously [64]. The DNA sequence of the clone was determined to verify its accuracy. Feeding RNAi was performed as described [65,64]. Briefly, bacteria were grown in liquid culture overnight and then plated on NGM plates containing 1 mM isopropyl-β-D-thiogalactopyranoside (IPTG). Fourth and fifth larval stage (L4) animals were transferred to plates with IPTG and 24 h later were transferred to fresh plates. Progeny were tested for engulfment or DTC migration defects.

In vitro binding. PCR products were made from the *abl-1* cDNA
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References


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