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that AXR4 may not be required for AUX1 trafficking in these cells. However, additional studies are required to confirm this possibility. These observations prompted us to investigate whether the trafficking of other membrane proteins is also dependent on AXR4 function. The auxin efflux facilitator PIN1 is normally found on the lower side of vascular cells (5) (Fig. 3D), whereas PIN2 is localized to the lower side of cortical cells and the upper side of epidermal cells (5) (Fig. 3E). We found that both PIN1 and PIN2 are localized normally in axr4 roots (Fig. 3, H and I). The localization of other plasma membrane proteins such as proton adenosine triphosphatase (PM H+-ATPase) is also normal in the axr4 mutant (Fig. 3, F and J). The axr4 mutation therefore appears to selectively disrupt trafficking of AUX1. We next investigated where AUX1 accumulates in the axr4 background by using a selection of endomembrane compartment markers (fig. S6). Pixel correlation analysis revealed that AUX1 overlaps most significantly with markers from the ER (fig. S6) like Sec12 (Fig. 3, K to M), suggesting that loss of AXR4 causes AUX1 to accumulate in the ER.

Recent work has demonstrated that for root gravitropism to occur, AUX1 must be expressed in both the LRC and expanding epidermal cells in order to transport gravity-induced lateral auxin gradients from gravisensing columella cells to graviresponsive epidermal cells (20). Localization results suggest that the gravitropic defect of axr4 seedlings is caused by the failure to traffic AUX1 to the plasma membrane in the epidermis (Fig. 3B). As a result, the axr4 mutation is expected to disrupt the AUX1dependent transfer of the lateral auxin gradient from the LRC to expanding epidermal cells. Consistent with this model, the auxin-responsive reporter IAA2::GUS is expressed normally in the LRC but is almost undetectable in the epidermis of the axr4 mutant (compare fig. S7, A and C). In contrast, IAA2::GUS expression is not detected in either the LRC or epidermal cells in the aux1 background (fig. S7B). The pattern of IAA2::GUS expression is consistent with the fact that AUX1 remains functional in the axr4 LRC but not in the epidermal cells. The axr4 mutant is resistant to 2,4-D, and because AUX1 may be functional in the LRC in axr4 mutants, this would suggest that functional AUX1 in the LRC alone is not sufficient for 2,4-D-sensitive root growth. We tested this possibility by transactivating AUX1 in the LRC or LRC plus epidermal cells (fig. S8) using Gal4 driver lines M0013 and J0951, respectively (20). When AUX1 was expressed in LRC alone using the GAL4 line M0013, the aux1 M0013 >> AUX1 line exhibited an auxinresistant root phenotype like that of axr4 (fig. S8, E and F) (20). However, when AUX1 was also expressed in the epidermal cells using the GAL4 line J0951, normal auxin response was restored (fig. S8, B and F). Our results suggest

that the *axr4* phenotype, including auxinresistant root growth and reduced gravitropism, is caused by defective AUX1 trafficking in epidermal cells.

AXR4 joins a growing list of ER accessory proteins that facilitate trafficking of plasma membrane proteins through the secretory pathway (21-23). In yeast cells, the Shr3 protein acts as a molecular chaperone of amino acid permeases, preventing their inappropriate aggregation in the ER membrane (21). Other structurally unrelated accessory proteins also appear to function as molecular chaperones of their cognate substrates (21). Although unrelated to Shr3p, AXR4 may have a similar function. Alternatively, because AXR4 is a member of the α/β hydrolase superfamily, it may facilitate AUX1 polar trafficking by posttranslationally modifying AUX1, causing it to be recognized as cargo destined for a particular plasma membrane face of the plant cell. However, validation of this or other mechanisms awaits further experimentation.

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Figs. S1 to S8 References

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CRACM1 Is a Plasma Membrane Protein Essential for Store-Operated Ca²⁺ Entry

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Store-operated Ca²⁺ entry is mediated by Ca²⁺ release–activated Ca²⁺ (CRAC) channels following Ca²⁺ release from intracellular stores. We performed a genome-wide RNA interference (RNAi) screen in *Drosophila* cells to identify proteins that inhibit store-operated Ca²⁺ influx. A secondary patch-clamp screen identified CRACM1 and CRACM2 (CRAC modulators 1 and 2) as modulators of *Drosophila* CRAC currents. We characterized the human ortholog of CRACM1, a plasma membrane–resident protein encoded by gene *FLJ14466*. Although overexpression of CRACM1 did not affect CRAC currents, RNAi-mediated knockdown disrupted its activation. CRACM1 could be the CRAC channel itself, a subunit of it, or a component of the CRAC signaling machinery.

 $R^{eceptor-mediated signaling in nonexcitable cells, immune cells in particular, involves an initial rise in intracellular Ca^{2+} due to release from the intracellular stores. The resulting depletion of$

the intracellular stores induces Ca^{2+} entry through the plasma membrane through CRAC channels (1-4). This phenomenon is central to many physiological processes such as T cell proliferation, gene transcription, and cytokine release (3, 5-7). Biophysically, CRAC currents have been well characterized (2, 8, 9), but the identity of the CRAC channel itself and the pathway resulting in its activation are still unknown. Recently, STIM1 (for stromal interaction molecule in Drosophila) was identified as an essential component of store-operated calcium entry (10, 11). This protein is located in intracellular compartments that likely represent parts of the endoplasmic reticulum (ER). It has a single transmembrane-spanning domain with a C-terminal Ca²⁺-binding motif that appears to be crucial for its hypothesized function as the ER sensor for luminal Ca²⁺ concentration. When stores become depleted, STIM1 redistributes into distinct structures (punctae) that move toward and accumulate underneath the plasma membrane. Whether or not STIM1 actually incorporates into the plasma membrane is controversial (10, 12, 13). Although STIM1 is required to activate CRAC currents, its presence or even its translocation appears not to be sufficient to mediate CRAC activation, because lymphocytes from patients with severe combined immunodeficiency (SCID) appear to have normal amounts of STIM1 levels and normal function, yet fail to activate CRAC channels (14). This suggests that other molecular components may participate in the store-operated Ca2+ entry mechanism.

To identify genes encoding the CRAC channel or other proteins involved in its regulation, we performed a high-throughput, genome-wide RNA interference (RNAi) screen in Drosophila S2R+ cells. The effect of knockdown of each of the ~23,000 genes was tested by fluorescence measurements of intracellular Ca²⁺ concentration in 384-well microplates with an automated fluorometric imaging plate reader (FLIPR, Molecular Devices). Changes in $[Ca^{2+}]_i$ were measured in response to the commonly used SERCA [sarcoplasmic and endoplasmic reticulum calcium adenosine triphosphatase (ATPase)] inhibitor thapsigargin, which causes depletion of Ca2+ from intracellular stores. An example of responses from this primary screen is illustrated in Fig. 1A, obtained from microplate no. 60. All 63 plates contained wells in which double-stranded RNA (dsRNA) against Rho1 served as negative control and dsRNA against stim1 as positive control. Higher resolution graphs of the realtime [Ca²⁺], imaging data are shown in Fig.

Fig. 1. Identification of CRACM1 and CRACM2 as crucial regulators of storeoperated Ca2+ entry in Drosophila. (A) Ca²⁺ signals measured in Drosophila S2R+ cells in the primary highthroughput screen using FLIPR. Representative FLIPR raw data file showing 384 minigraphs, each of which represents fluo-4 fluorescence change in an individual well with respect to time. Each plate contained the negative control dsRNA Rho1 in well A1 and the positive control dsRNA stim1 in well B1. (B) Fluo-4 fluorescence changes in relative fluorescence units (r.f.u.) obtained from cells treated with the indicated dsRNAs. Cells were kept in Ca2+-free solution and exposed to thapsigargin (2 μM), followed by addition of 2 mM Ca²⁺. The traces are representative of two independent repeats of the primary screen. (C) Same protocol as in (B) but for cells treated with CRACM2 dsRNA. (D) Normalized average time course of IP₃-induced (20 µM) I_{CRAC} measured in Drosophila



Kc cells. Currents of individual cells were measured at -80 mV, normalized by their respective cell size, averaged and plotted versus time (±SEM). Cytosolic calcium was clamped to 150 nM with 10 mM BAPTA and 4 mM CaCl₂. Traces correspond to untreated control [wild type (wt), black filled circles, n =10); Rho1 dsRNA (mock, open circles, n = 8); CRACM1 dsRNA (red circles, n = 6); and CRACM2 dsRNA (green circles, n = 9)]. (E) Averaged current-voltage (//V) data traces of I_{CRAC} extracted from representative cells at 60 s for currents evoked by 50-ms voltage ramps from -100 to +100 mV with leak currents subtracted and normalized to cell size (pF). Traces correspond to untreated control (wt, n = 9); CRACM1 dsRNA (n = 5); and CRACM2 dsRNA (n = 6).

1, B and C, from cells treated with dsRNA against Rho1 (mock) and stim1, as well as two genes we later identified as CRAC modulators 1 and 2 (CRACM1 and CRACM2). On the basis of inhibitory efficacy relative to positive and negative controls, we identified ~1500 genes that reduced Ca2+ influx to varying degrees (table S1). After eliminating numerous genes based on artifactual fluorescence signals or because they represent known housekeeping genes, cell cycle regulators, and so on, we eventually arrived at 27 candidate genes (table S2) that were subsequently evaluated in a secondary screen using single-cell patchclamp assays.

From the secondary patch-clamp screen, we identified two novel genes that are essential for CRAC channel function, CRACM1 (encoded by *olf186-F* in *Drosophila* and *FLJ14466* in human) and CRACM2 (encoded by *dpr3* in *Drosophila*, with no human ortholog). We measured CRAC currents in Drosophila Kc cells after inositol 1,4,5-trisphosphate (IP₃)-mediated depletion of Ca²⁺ from intracellular stores. Both untreated control wild-type cells and cells treated with an irrelevant dsRNA against Rhol (mock) responded by rapidly activating a Ca²⁺ current with the time course (Fig. 1D) and inwardly rectifying current-voltage (I/V) relation (Fig. 1E) typical of I_{CRAC} in mammalian (2) and Drosophila (15) cell types. In contrast, CRAC currents were essentially abolished in cells treated with dsRNA for CRACM1 and CRACM2. In some of the experiments on CRACM1, we also applied ionomycin (10 µM) extracellularly on top of the 20 µM IP₃ included in the patch pipette to ensure complete store depletion, but this also failed to induce I_{CRAC} (fig. S1C). Similarly, CRAC currents were also absent when passive store depletion was induced by the Ca²⁺ chelator

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BAPTA [(1,2-bis(*o*-aminophenoxy)ethane-*N*,*N*,*N*',*N*'-tetraacetic acid)] (fig. S1, A and B).

We studied the human ortholog of CRACM1, a 37.7-kD protein encoded by gene FLJ14466, to confirm that its function is conserved across species and that it is involved in store-operated Ca2+ entry. We used small interfering RNA (siRNA)-mediated silencing of human CRACM1 in human embryonic kidney cells (HEK293) and human T cells (Jurkat). The selective knockdown of CRACM1 message was confirmed by semiquantitative reverse transcription polymerase chain reaction (RT-PCR) analysis (Fig. 2A). Two different CRACM1-specific siRNA sequences caused a 60 to 70% inhibition of calcium influx in response to thapsigargin-induced store depletion in HEK293 cells (Fig. 2B). Patch-clamp recordings obtained from siRNA-treated cells, responding to intracellular IP3 perfusion, demonstrated a nearly complete inhibition of CRAC currents (Fig. 2, D and E). In Jurkat cells, siRNA-mediated inhibition of Ca2+ influx was close to 20% (Fig. 2C) and not as dramatic as in the HEK293 cells. However, $I_{\rm CRAC}$ in Jurkat cells was effectively reduced by both siRNA sequences (Fig. 2, F and G). The differences in the efficacy of suppressing the changes in [Ca2+], in HEK293 and Jurkat cells (see Fig. 2, B and C) are likely due to the different magnitudes of $I_{\rm CRAC}$ in these two cell types. CRAC current densities in HEK293 cells (~ 0.5 pA/pF) are much smaller than typically seen in Jurkat cells (~ 2.5 pA/pF), and a further inhibition may explain the more dramatic reduction in the Ca²⁺ signal than that observed in Jurkat cells (note that the remaining CRAC current densities in siRNA-treated Jurkat cells, while strongly reduced, are ~ 0.4 pA/pF and comparable to the normal CRAC current densities of untreated HEK293 cells). Taken together, these data indicate that CRACM1 is also a key modulator of storeoperated CRAC currents in human cells.

Given that the knockdown of CRACM1 inhibited CRAC activation, we wanted to know whether overexpression would enhance Ca²⁺ influx and CRAC current densities. HEK293, Jurkat, and RBL-2H3 cells were infected with a Myc-tagged CRACM1 and green fluorescent protein (GFP) retrovirus, and overexpression of the protein was confirmed in HEK293 cells by immunoprecipitation followed by Western blotting (Fig. 3A). However, we did not detect any increase in CRAC current amplitudes above control levels in either HEK293 (Fig. 3B) or Jurkat cells (fig. S2A) and only a slight increase in RBL cells (fig. S2B). These data suggest that CRACM1, although necessary for CRAC activation, does not in and of itself generate significantly larger CRAC currents.

An important question is whether CRACM1 localizes to the ER (as does STIM1) or to the



Fig. 2. Suppression of store-operated Ca²⁺ entry and I_{CRAC} by CRACM1 siRNA. (**A**) (Left) RT-PCR of CRACM1 mRNA from HEK293 cells infected with the indicated CRACM1-specific siRNAs and a scrambled sequence control. (Right) Control with primers specific for small ribosomal protein. (**B**) Fura 2–AM (pentaacetoxymethyl ester) fluorescence measurements of $[Ca^{2+}]_i$ in cells treated with scramble (control) or the two CRACM1-specific siRNAs in HEK293 cells. Cells were kept in Ca²⁺-free solution and exposed to thapsigargin (2 μ M), followed by addition of 2 mM Ca²⁺. The traces are representative of three independent experiments. (**C**) Same protocol as in (B), but for Jurkat cells. The traces are averages of three independent experiments. (**D**) Normalized average time course of IP₃-induced (20 μ M) I_{CRAC} measured in HEK293 cells treated with the indicated siRNAs (n = 9 to 13 for each group). [Ca²⁺]_i was clamped to near zero by 10 mM BAPTA. (**E**) Current-voltage (I/I/) data traces of I_{CRAC} from representative cells at 60 s for currents evoked by 50-ms voltage ramps from –100 to +100 mV in cells treated with the indicated siRNAs (n = 8 to 9).

plasma membrane. To address this question, we tagged CRACM1 on either end (Myc–C terminus and flag–N terminus) and transfected the constructs into HEK293 cells. After 24 hours, immunofluorescence confocal analysis revealed no staining in intact cells expressing either construct, which suggested that both tags are intracellular. After permeabilizing the cells, both constructs were detected by the fluorescent antibody and showed predominant peripheral staining of the plasma membrane (Fig. 3, C and D). These data fit well with the hydropathy profile of CRACM1, which predicts a topology of four transmembrane domains, with both ends facing the cytosol (fig. S2C).

In summary, our results demonstrate that the protein CRACM1 is essential for store-operated Ca^{2+} influx via CRAC channels. Although the overexpression of CRACM1 does not alter the magnitude of CRAC currents, the plasma membrane localization of this protein and the presence of multiple transmembrane domains point

Fig. 3. Overexpression of CRACM1. (A) Analysis of HEK293 cells for overexpression of CRACM1 by immunoprecipitation with antibodies against Myc or C-terminal His and immunoblotting with antibody against Myc. Control immunoprecipitation from empty vector-transfected cells did not show any bands. (B) Normalized average time course of IP₃-induced (20 µM) I_{CRAC} measured in HEK293 cells. Currents of individual cells were measured at -80 mV. normalized by their respective cell size, averaged, and plotted against time (± SEM). Cytosolic calcium was clamped to near zero by using 10 mM BAPTA. Traces correspond to cells transfected with GFP



alone (control, black circles, n = 13) and cells transfected with GFP plus CRACM1 (red circles, n = 14). (C) Immunofluorescence localization of CRACM1 in HEK293 cells visualized by confocal microscopy. Immunostaining for CRACM1–flag–N terminus (top) or CRACM1–Myc–C terminus (bottom) in intact (left) and permeabilized cells (right). (D) Same as bottom right panel of (C), but at higher magnification of selected cells to illustrate plasma membrane staining.

toward a direct role for CRACM1 in storeoperated calcium influx. A number of possible functions can be envisioned for CRACM1. First, CRACM1 could function as the CRAC channel itself. In this scenario, the unaltered CRAC currents in CRACM1 overexpressing cells might be due to a limiting factor upstream of CRAC channel activation (e.g., STIM1).

Second, CRACM1 could be a subunit of a multimeric channel complex, in which case the other subunit(s) could become the limiting factor(s) during overexpression. Finally, CRACM1 might function as a plasma membrane acceptor or docking protein, possibly for STIM1 or some other as-yet-unidentified component of the signaling machinery that ultimately leads to

CRAC channel activation and store-operated Ca^{2+} entry.

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Regulation of Adult Bone Mass by the Zinc Finger Adapter Protein Schnurri-3

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Genetic mutations that disrupt osteoblast function can result in skeletal dysmorphogenesis or, more rarely, in increased postnatal bone formation. Here we show that Schnurri-3 (Shn3), a mammalian homolog of the *Drosophila* zinc finger adapter protein Shn, is an essential regulator of adult bone formation. Mice lacking Shn3 display adult-onset osteosclerosis with increased bone mass due to augmented osteoblast activity. Shn3 was found to control protein levels of Runx2, the principal transcriptional regulator of osteoblast differentiation, by promoting its degradation through recruitment of the E3 ubiquitin ligase WWP1 to Runx2. By this means, Runx2-mediated extracellular matrix mineralization was antagonized, revealing an essential role for Shn3 as a central regulator of postnatal bone mass.

In contrast to their role in embryonic development, few genes are known to regulate osteoblast function during postnatal skeletal remodeling (1-3). The transcription factor Runx2 is an essential

component of skeletogenesis, as evidenced by the human autosomal dominant disease cleidocranial dysplasia, an inherited disorder of bone development characterized by clavicular hypoplasia and cranial and facial abnormalities, which is caused by Runx2 mutations (4–6). Runx2^{-/-} mice also exhibit a complete lack of both intramembranous and endochondral ossification due to the absence of osteoblasts, resulting in an unmineralized skeleton (6, 7). Runx2 is required for early commitment of mesenchymal stem cells into osteoprogenitors, and it also functions later in osteoblast differentiation to regulate the formation of the extracellular matrix (8).

Schnurri-3 (Shn3), a large zinc finger protein, was originally identified as a DNA

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