Suppression of arthritis-induced bone erosion by a CRAC channel antagonist

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ABSTRACT

Objective: We have shown in vitro and in vivo that osteoclast maturation requires calcium-release activated calcium (CRAC) channels. In inflammatory arthritis, osteoclasts mediate severe and debilitating bone erosion. In the current study, we assess the value of CRAC channels as a therapeutic target to suppress bone erosion in acute inflammatory arthritis.

Methods: Collagen-induced arthritis (CIA) was induced in mice. The CRAC channel inhibitor 3,4-dichloropropionaniline (DCPA) and a placebo was administered 1 day prior to collagen II booster to induce arthritis. Effects on swelling, inflammatory cell invasion in joints, serum cytokines and bone erosion were measured.

Results: Assays, by blinded observers, of arthritis severity showed that DCPA, 21 mg/kg/day, suppressed arthritis development over 3 weeks. Bone and cartilage damage in sections of animal feet was reduced approximately 50%; overall swelling of joints was reduced by a similar amount. Effects on bone density by µCT showed clear separation in DCPA-treated CIA animals from CIA without treatment, while differences between controls without CIA and CIA treated with DCPA differed by small amounts and in most cases were not statistically different. Response was not related to anticollegen titres. There were no adverse effects in the treated group on animal weight or activity, consistent with low toxicity. The effect was maximal 12–17 days after collagen booster, during the rapid appearance of arthritis in untreated CIA. At 20 days after treatment (day 40), differences in arthritis score were reduced and tumour necrosis factor α (TNF-α)-blocking molecules all of which can cause dangerous or debilitating side effects.

Conclusions: DCPA, a novel inhibitor of CRAC channels, suppresses bone erosion associated with acute arthritis in mice and might represent a new treatment modality for acute arthrits.

Rheumatoid arthritis (RA) has a chronic relapsing course with bouts of acute arthritis leading to destruction often of multiple joints and usually requiring lifelong therapy. While there are infectious causes of inflammatory arthritis and genetic predilections, typically RA also is idiopathic. Treatments for inflammatory arthritis include antimetabolites, steroids and tumour necrosis factor α (TNF-α)-blocking molecules all of which can cause dangerous or debilitating side effects.

We arrived at calcium-release activated calcium current (CRAC) inhibitors as a potential new means to treat acute arthritis through a series of studies of the role of calcium in osteoclast development. We found that the IP3R receptor calcium channel was necessary for osteoclast adhesion and activity, while others showed that IP3R-mediated release of calcium was required for activation of nuclear factor of activated T cells (NFATc1)—a critical signal for osteoclast formation. We then demonstrated that osteoclastogenesis was dependent on CRAC channels activated by release of calcium stores: suppression of CRAC via siRNA or inhibition by the CRAC antagonist, 3,4-dichloropropionaniline (DCPA) impaired differentiation of bone degrading osteoclasts. Further, CRAC is required for normal T-cell differentiation, also via NFATc1 activation. Thus a web of lymphocyte and macrophage differentiation events depend on CRAC, all of which might be involved in arthritis progression.
While total developmental suppression of CRAC causes severe bone deformities and skeletal defects. In adults, where skeletal development is mature, short-term treatment is unlikely to cause major skeletal effects. However, in long-term treatment immunological effects are likely. Moreover, since the Th17 response important, the pathogenesis of RA appears to be dependent on the CRAC channel Orai189 we reasoned that partial CRAC inhibition by DCPA might effectively blunt acute arthritis. Although several CRAC inhibitors are known, we pursued DCPA in part because of very low toxicity; exposure of humans to DCPA in mg/kg quantities has minimal apparent toxicity, although massive doses, approximately 4–5 g/kg, can cause fatal methemoglobinemia.

We performed a trial of suppression of acute arthritis using controlled-release DCPA, at 10.5 or 21 mg/kg/day, in mice with acute collagen-induced arthritis (CIA). DCPA caused a dramatic suppression of bone destruction and of articular inflammation at the higher dose. The studies used blinded observation of joint swelling over 5 weeks, and histological and μCT examination of sections of bone in the feet. In addition, we studied anticollagen antibodies and inflammatory cytokines, which did not correlate with arthritis response at the times studied. Osteoclast and T-cell numbers were also examined by immunohistochemistry in affected joints and while osteoclast numbers were reduced, T-cells were at similar density in inflammatory infiltrates in all groups.

MATERIALS AND METHODS

Animal studies

Six-week old male DBA/1 mice were purchased from Harlan Laboratories (Frederick MD) and housed in the WVU vivarium. These animals were acclimated at least 1 week prior to any experimental use. All animal procedures were approved by the WVU Institutional Animal Care and Use Committee.

Induction and assessment of arthritis

CIA was induced as described by Clutter et al. Briefly, bovine type II collagen (CII)(Elastin Products Co, Owensville, Missouri, USA) was emulsified in complete Freund’s adjuvant consisting of 10 mL of Freund’s Incomplete Adjuvant (Difco Laboratories, Detroit, Michigan, USA) plus 20 mg of Mycobacterium tuberculosis H37RA (Difco Laboratories). The CII (100 μg per animal; approximately 4 μg/kg) was injected intradermally on day 1 and 21 days later, a booster dose of 100 μg CII in Freund’s incomplete adjuvant (Difco Laboratories) was administered. Inflammation was apparent 4–8 days after the second dose, in 80% of treated joints. At day 20 after primary immunisation, time-release pellets (Innovative Research of America, Sarasota FL) containing DCPA or the placebo, calibrated to release the stated doses for 21 days, were placed subcutaneously. Power analysis indicated that at least eight animals per CIA group were required to provide a valid statistical sample. Since induction of CIA does not occur in 100% of the treated mice, 12 mice in each CIA-induction group were initially started in the experiment. Treatment doses included 0 mg/kg (placebo), 10.5 mg/kg/day of DCPA or 21 mg/kg/day of DCPA were compared. Four untreated controls, that is, no CIA or DCPA treatment, were also included.

Mice were monitored for arthritis and scored in a blinded manner as described by Clutter et al. Briefly, swelling of paws was be graded on scale from 0 to 4 indicating number of swollen digits. All paws were evaluated, so that the maximal arthritic index per mouse was 16. Additionally, hind paw swelling was measured using digital calipers on day 0, and each day on days 23–40. Analysis of the bones and joints for arthritis was performed on H&E stained sections of hind paws, by blinded observation. This scored synovial expansion and inflammation, joint damage including pannus and bone degradation, each on a scale of 0–3, with maximum score of 9. For histological analysis, two paws from each animal were analysed separately and blindly, and are calculated as two specimens per animal.

Serum analysis for antibodies and cytokines

Heart blood collected at the time of euthanasia on day 40 was used for analysis. Plasma was separated by centrifugation and frozen in aliquots at −20°C until used. Production of anti-CII antibodies was evaluated by ELISA (Rheumera, Astarte Biologics, Redmond, Washington, USA) and cytokine concentrations were measured using V-PLEX panels (Meso Scale Discovery, Rockville, Maryland, USA) using the methods prescribed by the respective manufacturers.

Antibody labelling of sections

Histological sections from the feet of animals euthanised at 40 days, were stained using standard immunohistochemical methods to measure the effect of DCPA on osteoclast bone interface and T-cell density. Osteoclast bone interface density was determined by anti-ATPα3 (TCIRG) labelling, and the effect on CD3 T-cell density was determined using anti-CD3 labelling. Anti-TCIRG1 quantification was mouse monoclonal (clone 6H3) antibody (Sigma-Aldrich) at 1:100 dilution and CD3 quantification used mouse monoclonal antibody anti-CD3 PC3/188A (raised against amino acids 156–168 of the cytoplasmic domain of human CD3ε) at a 1:100 dilution. Briefly, sections were blocked in phosphate-buffered saline (PBS) with 2% hydrogen peroxide for 5 min, then in PBS with 2% bovine serum albumin (BSA) for 2 h. The sections were incubated overnight with antibodies at indicated concentrations in PBS with 0.01% tween 20. After washing, sections were incubated for 1 h with biotinylated antimouse antibodies at 1:1000 dilution, washed again and incubated with streptavidin-horseradish peroxidase and diaminobenzidine substrate.
for 5 min. H&E counterstaining was performed to show tissue features. Imaging used a Nikon TE2000 inverted microscope, with 14-bit 2048×2048 pixel monochrome CCD camera and RGB filters to reconstruct colour (Spot, Sterling Heights, Michigan, USA).

**µCT and morphometry**

Analysis by µCT was as described. In brief, paws were scanned on a Viva CT40 instrument (Scanco, Bassersdorf, Switzerland) with three-dimensional (3D) reconstruction by the manufacturer’s instrument-specific software. The scan section increment was 20 µm; 3D reconstruction used a density cut-off of 211 mg/cm³. Analysis of trabecular thickness and trabecular spacing used the plate model.

**Statistics**

Continuous data obtained at one time point (day 40), such as anticollagen antibody titres, bone density, bone volume, concentration of cytokines, osteoclast area and per cent area of periosteal T cells, were analysed using analysis of variance (ANOVA) followed by Dunnett’s multiple comparison procedure (MCP) when the goal was to compare the CIA-treated groups to untreated controls. In addition, Tukey-Kramer analysis with adjustment for MCP was performed for continuous variables when the goal was to compare the differences among the CIA-treated groups. The hind paw swelling data, recorded over time (day 0 and 23–40), were analysed by the repeated measures ANOVA, using the autoregressive autocorrelation structure and followed by Tukey-Kramer analysis with adjustment for MCP. Data for anticollagen antibody titres and paw swelling were ln-transformed and reciprocal square-root transformed, respectively, before ANOVA due to their positive skewness. Data of categorical or ordinal nature, such as arthritis index, were analysed by non-parametric Kruskal-Wallis test for each day. In addition, arthritis index data were grouped into two categories based on the response, 0 for arthritis index equal to zero and 1 for all mice with arthritis index greater than 0. These data were also analysed by Kruskal-Wallis test for each day, followed by the non-parametric Wilcoxon rank sum multiple comparison. A 60% power of statistical test was based on the average arthritis index for each group and the expected SD. Data were analysed using JMP and SAS software (SAS Institute Inc, Cary, North Carolina, USA). Significance criterion α for all tests was 0.05.

**RESULTS**

In vivo analysis of CIA and the effect of DCPA treatment by continuous-release subcutaneous pellets. Mice showed no changes in body weight (data not shown), or increased lethargy or overt stress in any group, although severe inflammation of paws was induced by the CIA protocol. As measured by the arthritis index scoring each paw as 0–4 digits affected, on days 23–40 (after booster injections at day 20), 80% of mice responded to collagen injections by induction of arthritis (figure 1A). The collagen-treated animals receiving placebo (0 mg/kg/day DCPA) differed significantly from untreated controls, as expected (p<0.05 by Kruskal-Wallis test). The effect of 10.5 mg/kg/day of DCPA was modest and at most time points the results did not differ significantly by ANOVA from those of CIA animals treated with placebo (0 mg/kg/day DCPA). On the other hand, a dose of 21 mg/kg/day of DCPA reduced the arthritic index, strongly before day 37 and 20–50% at days 38–40. The arthritic index for animals treated with 21 mg/kg/day was statistically lower from that of CIA animals treated with placebo (Wilcoxon mean scores 14.5 vs 21, respectively, p<0.05 by Wilcoxon test, scores not shown). A scatter plot of the median arthritic scores of all animals on day 37 (day of maximum effect) is shown in figure 1B. In the CIA mice treated with 0 or 10.5 mg/kg/day DCPA >50% of the animals showed arthritis scores that variability was high. The majority of animals treated in the CIA animals treated with 21 mg/kg/day DCPA exhibited minimal arthritis scores, although three animals showed significant arthritis development. Hind paw thickness (figure 1C) was also measured with the average thickness for the untreated controls value of 2.29±0.03 mm. There were significant effect of treatment (p=0.02) and time (p<0.01) (data not shown) on paw thickness. Comparisons showed that on average, CIA-treated, low-dose DCPA (10.5 mg/kg/day) as well as placebo DCPA (0 mg/kg/day)-treated mice had significantly thicker paws (2.56±0.04 mm), than the untreated control mice (p<0.05 for each comparison). Those CIA-treated animals treated with 21 mg/kg/day DCPA had a paw thickness of 2.43±0.04 mm which was not significantly different from the untreated controls (p=0.52). There was a relatively high variability in the data which likely reflects the all-or-none nature of arthritis; in some of the few mice treated with 21 mg/kg/day DCPA that still developed arthritis, it was severe.

**Analysis of CIA in sections of feet**

Hind paws of animals were fixed, decalcified, paraffin embedded, cut and stained with H&E (figure 2A–C). Arthritis was scored by a blinded observer on a scale of 0–3 for inflammation in periarticular tissue with 3 being suppurative inflammation, 0–3 for involvement of cartilage and the joint space, with 3 requiring a pannus and 0–3 for degradation of periarticular bone, with 3 being full thickness destruction of the end plate. Representative examples of paws from an untreated animal showing no inflammation (figure 2A), a DCPA-treated CIA animal showing measurable but not severe arthritis (figure 2C), and a placebo-treated CIA animal with severe disease (figure 2B) are shown. The quantification of arthritis severity by a blinded observer is shown in figure 2D. The high-dose DCPA/CIA group was statistically different from the CIA/placebo group.
with \( p=0.024 \). The untreated group was statistically different from all other groups with \( p<0.01 \). Results for the low dose (10.5 mg/kg/day) treatment group were poorly separated from results for other CIA groups (the low-dose DCPA-treated animals were therefore omitted from further analyses). Figure 2E shows the anti-type II collagen (CII) titres. Mean anti-CII titres in the 0 mg/kg/day DCPA group were \( 7.1\pm2.1\times10^3 \), \( 5.1\pm1.1\times10^3 \) in the 10.5 mg/kg/day DCPA group and \( 8.3\pm2.8\times10^3 \) in the 21 mg/kg/day DCPA group; no significant differences among the CIA groups were found for anti-CII titres.

\( \mu \)CT

There were clear erosions in CIA mice which were not observed in high-dose DCPA-treated animals (figure 3A–C), in accord with histology (figure 2A–C); high-dose CIA/DCPA could not be distinguished visually from untreated controls. Low-dose CIA/DCPA (10.5 mg/kg/day) was not studied. Quantitative morphometry in the paws, the most severely affected site in the skeleton, showed that bone density and bone volume per total volume was significantly decreased in CIA/placebo mice. The CIA/high-dose DCPA treatment group had density indistinguishable from that of untreated controls (figure 3D) and showed a trend toward greater density than in CIA/placebo mice (\( p=0.08 \)). The bone volume/total volume for CIA/DCPA-treated mice did not differ from that of untreated controls, and it was statistically greater than for CIA/placebo animals (\( p=0.03 \)). Other parameters of bone morphometry including trabecular thickness and spacing were not different between groups (data not shown). This reflects the fact that the damage to the bone in arthritis is specific to the joints, which are excluded when determining bone trabecular parameters.

Effect of CIA with and without DCPA treatment on serum inflammatory cytokines

We determined the effect of DCPA treatment on key serum cytokines including TNF-\( \alpha \), IL-1 and IL-6. The results showed strong induction of the cytokines by CIA but, surprisingly, showed no effect of DCPA on cytokine production despite strong suppression of arthritis by DCPA (figures 1–3). Representative data from one of two separate analyses with essentially identical results is shown in figure 4. Interferon \( \gamma \) (IFN-\( \gamma \)), CXCL1, TNF-\( \alpha \), IL-1\( \beta \) and IL-6 (figure 4A–E) were all significantly increased in CIA/placebo animals compared to the untreated (no collagen injection) controls as expected. TNF-\( \alpha \) was also significantly increased in the CIA/high-dose DCPA mice compared to untreated controls. Comparison of CIA/placebo and CIA/high-dose DCPA groups revealed no significant reductions in any cytokines with CRAC inhibition. Based on these data, we hypothesise that the inhibitory effect of DCPA on osteoclasts was not secondary to changes in systemic cytokine levels, as these were not significantly affected. Nor did DCPA significantly affect production of IL-12p70 or

**Figure 1** The effect of 3,4-dichloropropionaniline (DCPA) treatment by continuous-release subcutaneous pellets on collagen-induced arthritis (CIA) as measured by the arthritic index. The data were gathered by a blinded observer. \( n=4 \) for untreated controls; in the CIA animals \( n=11 \) for animals treated with 0 mg/kg/day DCPA; \( n=10 \) for animals treated with 10.5 mg/kg/day DCPA; \( n=8 \) for animals treated with 21 mg/kg/day DCPA (green; closed circles) and untreated controls (red; closed square). There is a trend toward efficacy at intermediate times in CIA animals treated with 10.5 mg/kg/day DCPA (green; closed circles) but only the CIA animals treated with 21 mg/kg/day DCPA (yellow; closed triangles) had a durable effect. (B) Scatter plot showing the arthritis scores at day 37. This demonstrates that >50% animals in the CIA animals treated with 0 mg/kg/day DCPA (green; open circles) and 10.5 mg/kg/day DCPA (green; closed circles) show arthritis scores that exceed the mean and several animals showed the maximum arthritis score, whereas a majority of the animals /21 mg/kg/day DCPA (yellow; closed triangles) showed no or minimal arthritis scores. (C) Paw thickness, mean\( \pm \)SEM, for 4 control, 11 untreated CIA, and 8 high-dose DCPA/CIA animals. The difference between control and CIA animals is significant (*, \( p<0.05 \)); the high-dose DCPA animals are not different relative to the other groups (ns).
IL-10, which also were measured (not shown). IL-17 was also assayed but systemic levels were too low for meaningful comparisons (see Discussion).

Effect of CIA and DCPA on osteoclasts and T cells in tissue sections

Antibody labelling for TCIRG1/ATPa3 was used to determine the effect of the DCPA treatment on osteoclasts in tissue (figure 5A–D). TCIRG1 is the large subunit of the osteoclast-membrane hydrogen pump; it is not expressed by macrophages (nor in other cells in the marrow); in contrast, enzymes including cathepsin K and TRAP, which are highly expressed by osteoclasts and often used as osteoclast markers, are also expressed in macrophages and so are less specific.14 15 TCIRG1 labelling revealed a large and statistically significant reduction in the bone surface area covered by osteoclasts in CIA/DCPA-treated mice compared to CIA/placebo mice (figure 5D). In contrast, anti-CD3 labelling showed the density of T cells in inflammatory infiltrates to be similar in the CIA/placebo and CIA/DCPA groups (figure 5E).

Figure 2  The effect of 3,4-dichloropropionaniline (DCPA) treatment on collagen-induced arthritis (CIA) at 40 days. (A–C) The top frame shows an ankle bone in an untreated animal (untreated control, histological score 0). The middle frame shows severe arthritis in CIA treated with 0 mg/kg/day DCPA, with severe synovial inflammation, pannus in the joint space and severe bone degradation (histological damage score 8). The bottom frame shows the results from CIA animals treated with 21 mg/kg/day; there is inflammation, with synovial thickening, but it is not severe (histological damage score 4). All fields are 1.2 mm across. (D) Severity of arthritis in each group scored by a blinded observer based on sections as in A–C. The histological joint damage index (HJDI) in CIA animals treated with 21 mg/kg/day DCPA (yellow bar) is statistically different from those treated with 0 mg/kg/day (blue bar), p=0.024. The untreated group was different from all other groups, p<0.01. The CIA animals treated with 10.5 mg/kg/day (green bar) were poorly separated on day 40 HJDI, and were not studied further. Both hind feet of each animal were individually processed and blindly scored, so the number of measurements is twice the number of animals and expressed as the mean±SEM HJDI. Animal numbers are as stated for figure 1. (E) Anti-type II collagen titre in CIA mice. The anti-type II collagen titre in CIA animals treated with 21 mg/kg/day DCPA (yellow bar) was 8.3 (±2.8)×10³ and was not statistically different from those treated with 0 mg/kg/day (mean titre=7.1 (±2.1) × 10³; blue bar). The mean anti-type II collagen titre of CIA animals treated with 10.5 mg/kg/day (green bar) was 5.1 (±1.1) × 10³ and also was not statistically different from the other CIA animals. The anti-type II collagen titres in the untreated group were all below the level of detection. Values shown are the respective group mean±SEM.
complement therapy that targets the in
serum cytokines, suggesting that the approach may
These effects were independent from suppression of
past days, when arthritis effect diminished (figure 1A).
ment where it interacts with Orai1.18 Using
electrophysiological CRAC measurement and fluores-
cence analysis of STIM1/Orai1 association, we deter-
mixed that DCPA inhibits CRAC channel inhibition by
destabilising STIM1/Orai1 interaction, leading to CRAC
channel closure.4 In that study, we further showed that
DCPA inhibits osteoclast formation in vitro; we now show
that DCPA-induced CRAC channel inhibition leads to
inhibition of osteoclast formation and function in vivo
in an arthritis model.
Human Orai1 defects are now known, and are asso-
ciated with defects including hypocalcaemia and mus-
cular abnormalities.19 Although our mouse model
showed no signs of serious metabolic or muscular
anomaly, in larger animal studies it will be important
to monitor these parameters. Oddly, some Orai1 poly-
morphisms also are associated with increased serum
calcium and kidney disease, although the mechanisms
are unclear.20 A dysfunctional STIM1 mutation, that is,
in the calcium sensor required to activate Orai1 com-
plexes, causes a syndrome with multiple developmental
anomalies, in larger animal studies it will be important
complete absence of CRAC is probably lethal during
earliest development or soon after birth. However, after
normal development, at least for a time frame of
3–4 weeks, suppression of CRAC appears to be rela-
tively benign.
Analysis of inflammatory cytokines showed that DCPA
treatment at 21 mg/kg/day did not affect TNF-α, IL-1,
IL-6, CXCL1, and IFN-γ (figure 4) all of which were
upregulated 2.5-fold to >10-fold by the CIA. We did not
detect significant IL-17; other work has shown that IL-17
after adjuvant-induced arthritis, at 35 days, had serum
concentrations <1 ng/mL. 35 days after arthritis indu-
ction, and which varied only approximately 20% between
untreated animals and arthritic animals,24 so it is likely
our assay was insufficiently sensitive to determine IL-17
effects. There is strong evidence for the involvement of
IL-17 in cellular signalling involved in induction of
Figure 3 μCT showing loss of bone in untreated animals
and effect of high-dose 3,4-dichloropropionaniline (DCPA).
(A–C) Examples of forelimb metacarpals in untreated mice,
CIA mice treated with 0 mg/kg/day and CIA mice treated with
21 mg/kg/day DCPA after euthanasia at day 40. Arrows in (B)
show cortical bone defects. (D–E) Bone density in total
volume of bone and bone volume per total volume in four
untreated controls and CIA animals treated with 0 (blue bar;
=9) or 21 (yellow bar; =9) mg/kg/day DCPA each. For both
measures the CIA animals treated with 0 mg/kg/day DCPA
has significantly reduced bone; the CIA animals treated with
21 mg/kg/day DCPA is not different from the untreated control
but is statistically greater than CIA animals treated with 0 mg/
kg/day in BV/V.

DISCUSSION
We report an in vivo trial of a CRAC inhibitor on
inflammatory arthritis using the mouse CIA model.12
The theoretical basis for the observed effect is that
CRAC currents are essential to normal osteclast de-
velopment, and also are involved in T-cell activation which
also promotes arthritis.4 5 Our results include a highly
significant decrease in joint swelling, inflammatory infil-
trate and bone damage. Despite the manufacturer’s
statement that DCPA release is constant, it is likely that
the amount of released DCPA decreased during the
past days, when arthritis effect diminished (figure 1A).
These effects were independent from suppression of
serum cytokines, suggesting that the approach may
complement therapy that targets the inflammatory cyto-
kines. The effects on many bone density and trabecular
bone parameters showed uniformly that DCPA reduced
the effects of CIA on bone health, although there was
variability in statistical significance between individual
parameters. On the other hand, an important unre-
solved issue is that it is likely, indeed almost certain,
arthritis, but it is not a major secreted serum protein; indeed, some amount of Orai1 signalling is believed to be required to support TH17 cell development. Typically, IL-17 production is assayed in tissue sections or cells, and further work will be needed to resolve the contribution of IL-17 suppression to the reduction of arthritis severity.

Recently, it was reported that another inhibitor of store-operated calcium entry, 4-methy-4′-[3,5-bis (trifluoromethyl)-1H-pyrazol-1-yl]-1,2,3-thiadiazole-5-carboxanilide, also called YM-58483 or BTP-2, at 10 mg/kg suppressed CIA and suppressed inflammatory cytokines IL-1, IL-6 and TNF-α to near-baseline levels at 10 days after induction. While it is expected that other inhibitors of CRAC channels would function similarly in suppressing arthritis, the differences in response of inflammatory cytokines are very interesting, although not unexpected. The inhibitor BTP2 suppresses cytokines including IL-2 and calcium influx by other calcium channels, specifically TRPM4, at low nanomolar concentrations. Overall, it is likely that BTP2 effects overlap those of DCPA, but that BTP2 has a range of effects that may be, at least in part, unrelated to CRAC suppression. It will be important to test additional time points to determine if DCPA has effects on some cytokines that were not resolved, but, importantly, at a time when skeletal effects were clearly suppressed by DCPA, IL-1, IL-6 and TNF-α were essentially identical in CIA with and without DCPA treatment (figure 4).

The clear effect of DCPA treatment on arthritis, despite no change in serum inflammatory factors, suggests that the direct blockade of CRAC activity by DCPA mediates the reduction in osteoclast maturation. This model (figure 6) will require further testing and refinement. However, the independence of the DCPA effect from suppression of the assayed serum inflammatory cytokines suggests that Orai1 antagonists may be valuable modalities for treating inflammatory arthritis, and possibly RA, by a mechanism independent of inflammatory cytokine suppression. However, serum cytokine levels may not reflect the levels of these cytokines within the joint spaces. It is likely that long-term strong

Figure 4  Effect of CIA and treatment with 3,4-dichloropropionaniline (DCPA) on production of key serum inflammatory factors. (A) IFN-γ. (B) CXCL1. (C) TNF-α. (D) IL-1β. (E) IL-6. One of two multiplex assays with similar results is shown. All of the cytokines shown were induced strongly with the exception of IL-10 (not shown). No cytokines differed between CIA animals treated with 0 (blue bar) or 21 mg/kg/day (yellow bar) DCPA. Statistical significance between groups is denoted by lower case letters above the bar. Mean levels not connected by the same letter are significantly different at p<0.05. IFN-γ, Interferon γ; IL-6, TNF-α, tumour necrosis factor α.

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Inhibition of CRAC would result in severe immunological or bone effects, based on the knockout mouse phenotype, but treatment for several weeks had no effect on animal health, activity, or weight.

In conclusion, treatment of CIA with 21 mg/kg/day of DCPA dramatically suppressed bone erosion and bone loss by μCT that is associated with CIA, with no apparent effect on the general health of the animals or the increase in serum inflammatory growth factors characteristic of acute arthritis. This remarkable ability of a CRAC channel inhibitor to regulate bone erosion without interfering with immune function may lead to a valuable new tool for the treatment of inflammatory arthritis.

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