Abstract

Inflammatory bowel disease (IBD) encompasses a range of intestinal pathologies, the most common of which are ulcerative colitis (UC) and Crohn's Disease (CD). Both UC and CD, when present in the colon, generate a similar symptom profile which can include diarrhea, rectal bleeding, abdominal pain, and weight loss. Although the pathogenesis of IBD remains unknown, it is described as a multifactorial disease that involves both genetic and environmental components.

There are numerous and variable animal models of colonic inflammation that resemble several features of IBD. Animal models of colitis range from those arising spontaneously in susceptible strains of certain species to those requiring administration of specific concentrations of colitis-inducing chemicals, such as dextran sulphate sodium (DSS). Chemical-induced models of gut inflammation are the most commonly used and best described models of IBD. Administration of DSS in drinking water produces acute or chronic colitis depending on the administration protocol. Animals given DSS exhibit weight loss and signs of loose stool or diarrhea, sometimes with evidence of rectal bleeding. Here, we describe the methods by which colitis development and the resulting inflammatory response can be characterized following administration of DSS. These methods include histological analysis of hematoxylin/eosin stained colon sections, measurement of pro-inflammatory cytokines, and determination of myeloperoxidase (MPO) activity, which can be used as a surrogate marker of inflammation.

The extent of the inflammatory response in disease state can be assessed by the presence of clinical symptoms or by alteration in histology in mucosal tissue. Colonic histological damage is assessed by using a scoring system that considers loss of crypt architecture, inflammatory cell infiltration, muscle thickening, goblet cell depletion, and crypt abscess. Quantitatively, levels of pro-inflammatory cytokines with acute inflammatory properties, such as interleukin (IL)-1β, IL-6 and tumour necrosis factor (TNF)-α, can be determined using conventional ELISA methods. In addition, MPO activity can be measured using a colorimetric assay and used as an index of inflammation.

In experimental colitis, disease severity is often correlated with an increase in MPO activity and higher levels of pro-inflammatory cytokines. Colitis severity and inflammation-associated damage can be assessed by examining stool consistency and bleeding, in addition to assessing the histopathological state of the intestine using hematoxylin/eosin stained colonic tissue sections. Colonic tissue fragments can be used to determine MPO activity and cytokine production. Taken together, these measures can be used to evaluate the intestinal inflammatory response in animal models of experimental colitis.

Video Link

The video component of this article can be found at http://www.jove.com/video/3678/.

Protocol

1. Murine model of DSS-induced acute colitis

1. Add dextran sulphate sodium (DSS) to autoclaved drinking water to the desired final concentration (1-5%) (wt/vol) (i.e. To make a 5% DSS solution, add 25 g of DSS powder to 500 mL of autoclaved water). Stock solution can be left at room temperature for up to one week or at 4°C until use.

2. In a biosafety hood, pour stock DSS solution into 50 mL Falcon tubes (one needed per cage). Keep the stock solution to refill tubes when needed.

3. Replace drinking water in each mouse cage with the DSS solution (in the 50 mL Falcon tubes) (The duration will depend on the DSS regimen that is used. For example, using 6-8 week old male C57BL/6 mice, we administer a 5% DSS solution for a total of five days). Mice should not have access to any other water source. Control mice are given autoclaved drinking water without DSS.

4. Weigh mice daily and record the amount of DSS consumed per day. Top each bottle to 50 mL after recording DSS levels. This is to measure the approximate volume of DSS consumed per cage per mouse throughout the duration of the experiment. In our studies, we use a 5% DSS solution for 5 days with male C57BL/6 mice. Significant weight loss, altered stool consistency and signs of fecal blood are seen as early as day 3 using this particular DSS regimen. During the DSS administration, mice exhibit pronounced weight loss (around 5-10% of their initial weight on day 5) with weight loss greater than 20% of initial weight with dehydration and diarrhea to be the significant physiological indicator that an animal is at or near endpoint. If the animal is given access to normal water after the 5 days of 5% DSS, it will recover within 7 days. All experiments should be approved by the institution's animal ethics committee and be in accordance with the approved Animal Utilization Protocol (AUP).
5. During the duration of the experiment, a disease activity index (DAI) score can be assessed to evaluate the clinical progression of colitis. The DAI is the combined score of weight loss compared to initial weight, stool consistency, and bleeding. Scores are defined as follows: weight loss: 0 (no loss), 1 (1-5%), 2 (5-10%), 3 (10-20%), and 4 (>20%); stool consistency: 0 (normal), 2 (loose stool), and 4 (diarrhea); and bleeding: 0 (no blood), 1 (Hemoccult positive), 2 (Hemoccult positive and visual pellet bleeding), and 4 (gross bleeding, blood around anus). DAI can be scored daily during the duration of the DSS treatment.

6. At the time point of choice, weigh and sacrifice mice. Mice can be euthanized by cervical dislocation following inhalation of isoflurane or by another method approved by the institution's animal facility.

2. Collect colonic tissue samples

1. Expose the ventral side of the animal and wet the abdomen area with a 70% ethanol solution. At this point, refer to Table 1 and make note of any signs of rectal bleeding (blood present at the anal orifice) or rectal prolapse in each animal.

2. Use standard dissecting scissors to incise the abdomen by making a ventral midline incision.

3. Locate the colon and transect the colon as close to the colorectal margin as possible to free the distal colon.

4. Carefully and slowly pull out the whole colon, detaching it from the surrounding mesentery.

5. Transect the colon at the colonocecal margin to free the proximal end of the colon. Feces can be removed by rinsing colon with sterile PBS by using a gavage needle attached to a 3 or 5 mL syringe or by carefully squeezing it out using a pair of bent tweezers/forceps. Using the whole colon, assess for damage (See Section 3.1)

6. Tissue sampling for histological analysis and other assays can be done by cutting 0.5 cm to 1.0 cm long colonic fragments making note of which area the sample is from (i.e. proximal, middle, or distal).

7. Tissue samples to be used for assays can be individually placed in 1.5 mL eppendorf tubes and frozen in liquid nitrogen and stored until use at -70°C.

3. Assessment of colitis severity

1. Macroscopic or disease severity score is assessed terminally by an unbiased observer using a previously published scoring system (Table 1).

   8 Stool consistency can be assessed by using a pair of forceps and pressing down on the feces to determine consistency. To determine a score for blood in the feces, note the colour of the feces (i.e. black stool versus light brown stool) and further validate using a Hemoccult test. Using the scoring system, determine a score for each of the conditions. The final macroscopic score for each animal is the sum of each individual score.

2. To evaluate histological damage of colitis severity, cut a small fragment (0.5 cm) of the colon, place in a tissue cassette and submerge in buffered 10% formalin solution. Prepare 5 μm paraffin embedded cross sections and stain sections with hematoxylin/eoxin (H&E) using the appropriate procedures. Colon fragments can be taken from the proximal, mid-colon, or distal section of the colon.

3. H&E stained colonic tissue sections are scored by a blinded observer using a previously published system for the following measures: crypt architecture (normal, 0 - severe crypt distortion with loss of entire crypts), degree of inflammatory cell infiltration (normal, 0 - dense inflammatory infiltrate), 3), muscle thickening (base of crypt sits on the muscularis mucosae, 0 - marked muscle thickening present, 3), goblet cell depletion (absent, 0- present, 1) and crypt abscess (absent, 0- present, 1). The histological damage score is the sum of each individual score. It should be noted that unlike human UC, crypt abscesses are not characteristic of this model and are rarely seen; microscopic ulcerations are also rare. If multiple colon sections were stained, histological scores between similar sections should be used to determine the final score for each area (i.e. histological score in proximal colon versus histological score in distal colon).

4. Prepare stock solutions of reagents for assays

1. Prepare a 50 mM solution of Potassium Phosphate buffer by adding solution B (K_2HPO_4, 8.7 g of dibasic potassium phosphate in 1 L of dH_2O) to solution A (KH_2PO_4, 6.8 g of monobasic potassium phosphate in 1L of dH_2O) until a pH of 6.0 is achieved. Remaining solutions can be stored in fridge (2-8°C) until future use.

2. Prepare hexadecyltrimethylammonium bromide (HTAB) buffer by adding 5 g HTAB into 1 L of Potassium Phosphate buffer (50 mM, pH=6.0). Gently heat to dissolve and store at 2-8°C until use. When required, heat to re-dissolve.

3. Prepare lysis buffer for tissue homogenization for protein analysis by adding 10 mL of 1M tris-hydrochloric acid (pH=8.0), 6 mL of 5M sodium chloride and 2 mL of Triton X-100 to 182 mL of sterilized distilled water. Triton X-100 is very viscous at room temperature and thus, should be gently warmed prior to use. The prepared lysis buffer can be stored at -20°C until use.

5. Sample preparation for assays

1. Sample preparation for MPO analysis.

   1. Remove samples from -70°C and place on ice. Record the weight of each sample after removing any visible feces or fat by using a bent forcep/tweezer and place into a 2mL Eppendorf Safe-Lock microcentrifuge tube (or any tube that can be used with a homogenizer). Samples should be kept on ice at all times. It is important to note that similar colon fragments should be used from each biological replicate (i.e. distal sections only or proximal sections only).

   2. Add homogenizer bead to each sample tube.

   3. Add the appropriate amount of HTAB buffer according to tissue weight. If tissue weight is less than 25 mg, add the buffer at a ratio of 12.5mg/mL; if tissue weight is between 25-50, add at a ratio of 25mg/mL. If tissue weight is greater than 50, add buffer at a ratio of 50mg/mL.

   4. Homogenize with a tissue homogenizer for 4 min at 30 Hz. Repeat if tissue is not fully homogenized.

   5. Remove homogenizer bead and centrifuge solution for 6 min (13400 x g, 4°C).

   6. Collect supernatant and discard the resulting pellet. Supernatent can be stored at -70°C until use.

2. Sample preparation for cytokine analysis.

   1. Repeat steps 5.1.1. to 5.1.2.

   2. Add 50 μl of protease inhibitor cocktail (PIC) to 10 mL of prepared lysis buffer.

   3. Add 1 mL of the PIC and lysis buffer solution to each sample irrespective of weight.

   4. Homogenize for 5 min at 30 Hz. Repeat if tissue is not fully homogenized.

   5. Remove homogenizer bead and centrifuge solution for 5 min at 3300 x g.
6. Quantification of inflammatory markers

1. MPO activity assay
   1. Prepare o-dianisidine dihydrochloride (o-dianisidine) solution by combining 16.7 mg of o-dianisidine dihydrochloride, 90 mL of dH₂O, and 10 mL of potassium phosphate buffer. This solution should be prepared fresh for every assay.
   2. Add 7 μL of tissue homogenate (prepared in section 5.1) in triplicate into a 96-well plate.
   3. Add 50 μL of diluted H₂O₂ (4 μL of 30% H₂O₂ diluted in 96 μL of dH₂O) to the o-dianisidine mixture.
   4. Use a multi-channel pipette to add 200 μL of the o-dianisidine mixture containing H₂O₂ to each of the wells.
   5. Measure absorbance at 450 nm using a spectrophotometer. Take three readings at 30 second intervals.
   6. Calculate MPO activity. MPO activity is measured in units (U) of MPO/mg tissue, where one unit of MPO is defined as the amount needed to degrade 1 μmol of H₂O₂ per minute at room temperature. Considering that one unit (U) of MPO= 1 μmol of H₂O₂ split and that 1 μmol of H₂O₂ gives a change of absorbance of 1.13 x 10⁻² nm/min, units of MPO in each sample is determined as change in absorbance ([ΔA(t₂-t₁)]/Δmin x (1.13 x 10⁻²)). To get units per mg of tissue, use the tissue: buffer ratio. For example, if a tissue: buffer ratio of 50 mg/mL was used, in 7 μL of homogenate, there is 0.35 mg of tissue. Therefore, to get units per mg tissue, divide the units of MPO by 0.35. A sample calculation using absorbance values (nm) is included below (assuming that the sample has been added in triplicate):

<table>
<thead>
<tr>
<th>Sample</th>
<th>Time 0 sec</th>
<th>Time 30 sec</th>
<th>Time 60 sec</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>0.048</td>
<td>0.061</td>
<td>0.074</td>
</tr>
<tr>
<td>2</td>
<td>0.048</td>
<td>0.061</td>
<td>0.073</td>
</tr>
<tr>
<td>3</td>
<td>0.051</td>
<td>0.065</td>
<td>0.078</td>
</tr>
</tbody>
</table>

i. Average at Time 0 sec= (0.048 + 0.048 + 0.051)/3= 0.049 nm
ii. Average at Time 30 sec= 0.0623 nm
iii. Average at Time 60 sec= 0.075 nm
iv. Change in absorbance (ΔA) from 0 to 30 sec/ mg tissue (assuming 50 mg/mL of tissue: buffer ratio)= [(0.0623-0.049)/(1.13 x 10⁻²)]/0.35= 3.363
v. Change in absorbance (ΔA) from 30 to 60 sec/ mg tissue= 3.211
vi. *MPO activity (U/mg tissue)= average of ΔA(0-30) and ΔA(30-60)= 3.287

2. Quantification of pro-inflammatory cytokines by ELISA
   1. Cytokine (IL-1β, IL-6 and TNF-α) levels are determined using commercially available enzyme-linked immunosorbent assay (ELISA) kit (Quantikine Murine; R&D Systems).
   2. Absorbance values from each ELISA is normalized using a Bradford protein assay respective to each sample and is expressed in units of pg/mg of protein.

7. Representative Results

Administration of the appropriate DSS regimen will induce acute colitis in mice. During the duration of the DSS treatment, DAI can be used to assess and evaluate the clinical progression of disease. Animals treated with DSS will show significant weight loss compared to their initial weights, loose stools and fecal bleeding (Figure 1). Upon sacrifice and examination of the colon, the severity of colitis is macroscopically scored based on shortening of colon length, colonic bleeding, fecal bleeding, loosening of stool consistency, and signs of rectal bleeding compared to controls treated with water only (Figure 2 & Table 1). Cross sections of colonic tissue samples stained with H&E will have higher histological scores for DSS-treated colons versus water-treated controls (Figure 3). To further characterize the extent of inflammation in DSS-treated mice, MPO activity can be assessed from homogenized colonic tissue samples. DSS-treated colons will have higher MPO activity compared to controls (Figure 4). In addition, this is associated with increased levels of pro-inflammatory cytokines (IL-1β, IL-6, TNF-α) (Figure 5).

Figure 1. Male, C57BL/6 mice were given 5% DSS in drinking water for 5 days. DAI scores were assessed daily for each animal and were averaged per day for each group (mean ± SEM, n=4 mice/group).
Figure 2. C57BL/6 mice were given 5% DSS in drinking water for 5 days. Control mice received water without DSS. Macroscopic damage score/disease severity scores were blindly assessed on day 5 post DSS-induced colitis. Colonos isolated from mice that received DSS have higher macroscopic damage scores (rectal bleeding, rectal prolapse, diarrhea, colonic bleeding) indicating greater disease severity (mean ± SEM, n=4 mice/group).

Figure 3. C57BL/6 mice were given 5% DSS solution in drinking water in order to induce colitis. Control mice received water without DSS. (A) Histological scores were blindly scored using H&E stained colonic tissue sections collected on day 5 post-DSS administration. (B) DSS-treated samples show more histological damage (more cellular infiltration, more goblet cell depletion, greater distortion/damage to crypt architecture) compared to (C) controls (mean ± SEM, n=4 mice/group). In (B) and (C), asterisk (*) indicates area of goblet cell depletion and distortion of crypt architecture; number sign (#) indicates cellular infiltration.

Figure 4. All mice were sacrificed on day 5 post administration of DSS and colonic tissue samples were collected to assess MPO activity. Severity of DSS induced colitis is associated with higher levels of MPO activity compared to controls (mean ± SEM, n=4 mice/group).
Figure 5. In addition to higher MPO levels, severity of DSS-induced colitis is also associated with an increased level of pro-inflammatory cytokines such as IL-1β, IL-6, TNF-α (mean ± SEM, n=4 mice/group).

Table 1. Macroscopic/Disease Severity Score

<table>
<thead>
<tr>
<th>Score</th>
<th>Rectal bleeding</th>
<th>Rectal Prolapse</th>
<th>Stool Consistency</th>
<th>Blood</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>None</td>
<td>None</td>
<td>Normal</td>
<td>Normal</td>
</tr>
<tr>
<td>1</td>
<td>Red</td>
<td>Signs of prolapse</td>
<td>Soft</td>
<td>Red</td>
</tr>
<tr>
<td>2</td>
<td>Dark Red</td>
<td>Clear prolapse</td>
<td>Very Soft</td>
<td>Dark red</td>
</tr>
<tr>
<td>3</td>
<td>Gross Bleeding</td>
<td>Extensive prolapse</td>
<td>Diarrhea</td>
<td>Black</td>
</tr>
</tbody>
</table>

Discussion

DSS colitis is a widely used chemically induced model of intestinal inflammation. In this model, mice are given drinking water supplemented with DSS, which is thought to be toxic to gut epithelial cells and disrupt the integrity of the mucosal barrier. Administration of DSS induces an acute colitis that is characterized by loose stool, fecal bleeding, and infiltration with granulocytes. During DSS administration, colitis is usually associated with significant weight loss and presence of blood in the stool, which can be assessed by a fecal occult blood test. After collecting colonic tissue samples, colitis severity can be characterized by macroscopic examination of the colon and histological analysis of H&E stained colonic cross-sections using previously established scoring systems.

When following this protocol, it is important to note that the severity of colitis induced by DSS is species and strain specific. In addition, differences in the intestinal microflora between different animal facilities and housing rooms may alter the outcome of DSS administration. Thus, initial studies using DSS may be required to optimize the dosage and duration of DSS treatment. Failure to optimize these variables may result in a high incidence of death or little to no colitis. Once optimized, this model can be used as a highly reproducible model of colitis with a low rate of mortality. In addition, it is important to use DSS of the specified molecular weight. Other forms of DSS salt reagent may not produce colitis or may lead to high incidence of death.

Induction of colitis using this model results in severe macroscopic and histological damage, which is associated with increased MPO activity as well as higher levels of pro-inflammatory cytokine production. Aggranulocytes, such as lymphocytes and monocytes, are important sources of pro-inflammatory cytokines, whereas MPO is an enzyme contained within granulocytes such as neutrophils (and to a lesser extent monocytes and macrophages). This protocol can also be used to measure MPO activity in colonic tissue samples treated with dinitrobenzene sulfonic acid (DNBS)-induced colitis. DNBS-induced colitis is a well-characterized T-cell mediated transmural inflammation of the colon that is administered by intrarectal instillation of the DNBS substance in ethanol. Ethanol is used to break down the mucosal barrier, allowing for the DNBS to haptenize to autologous or microbiota proteins stimulating a host immune response to the hapten-modified self-antigens. One advantage of this model over the DSS-induced model is that the eliciting agent is known. The mechanism by which DSS initiates colitis, however, remains to be determined. Interestingly, studies have shown that DSS administration causes colitis in natural killer cell-deficient, and T- and B-cell deficient mice, suggesting that these cells (associated with the adaptive immune system) may not be critical in the induction of colitis and thus, this model may be suitable for studying the role of the innate immune system in the generation of colitis.

For the most accurate measure of MPO activity, we have found that regardless of which colitis model is used, MPO levels should be determined within the first week of tissue collection as MPO activity tends to decrease over time. MPO activity can be used as a surrogate marker of inflammation. However, quantification of tissue cytokines and histological scoring are integral to facilitate the complete assessment of the inflammatory response during colitis.

Disclosures

No conflicts of interest declared.

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References


