# **Acceptance Program Requirements**



# Cleansers for Removable Prostheses

ADA American
Dental
Association®
Council on
Scientific Affairs



# **Acceptance Program Requirements**

This document outlines specific category requirements. Please also refer to the General Guidelines for Participation in the ADA Seal of Acceptance Program.

Category: Cleaners for Removable Prostheses

Purpose: The Acceptance Program applies to over-the-counter dental products for which

safety and efficacy has been established by laboratory and/or clinical evaluations where appropriate. Accordingly, the purpose of these requirements is to provide a structure upon which cleaners for removal prosthesis can be considered for ADA

Acceptance.

**Scope:** These requirements apply to materials and agents used to remove soft debris from

removable prosthesis, which include full and partial dentures with acrylic resin bases and partial dentures with acrylic resin bases and metal frameworks and/or clasps; to remove stains from removable prostheses; and to eliminate unpleasant

odors from these prostheses.

Notice Regarding Submission of Copyrighted Materials: To make the review of submissions to the ADA Acceptance Program as efficient as possible, the Council on Scientific Affairs provides copies of submitted materials to Council members and consultant reviewers, and also posts submitted materials to an area of the ADA's web site the access to which is restricted to Council members and staff.

By making a submission, you are representing and warranting to the Council on Scientific Affairs and the ADA that you have obtained sufficient permission(s) from the copyright owner(s) of any copyrighted material included with your submission to allow for the publication and distribution of that material by the ADA as described above, and agree to indemnify and hold ADA harmless from any and all claims arising from such publication or distribution.

Questions can be directed to adaseal@ada.org.



# 1. SEAL STATEMENT

The following statement applies to products approved under the below-listed criteria:

"The ADA Council on Scientific Affairs' Acceptance of (Product Name) is based on its finding that the product is safe and has shown efficacy cleaning removable prostheses, when used as directed."

Format for product packaging:

• Helps clean removable prostheses

# 2. SUBMISSION DIRECTIONS

- **A.** Submissions are to be sent in electronic format (email) to <a href="mailto:adaseal@ada.org">adaseal@ada.org</a>. Additional instructions will be provided regarding shipment of necessary samples.
- **B.** The submission fee is a one-time, non-refundable fee and is required before review begins. Maintenance fees are billed to the company in January of every year.
- C. The review timeline for new submissions is typically 4-6 weeks after all materials have been received. The decision to award the ADA Seal to a new product is made by the Council on Scientific Affairs. Family submissions may take anywhere from 2-4 weeks to review. Eligibility criteria for Family Submissions are outlined in the Guidelines for Participation in the ADA Seal of Acceptance Program.

Note: This is an estimated timeline. Extended review time may be required if additional information or clarification is needed from the manufacturer.

- D. When a product is classified as "Accepted" and is awarded the ADA Seal of Acceptance, the Acceptance period is five years. Manufacturers will be contacted approximately six months before the expiration of the current Acceptance period to complete the requirements for the next five-year Acceptance period.
- **E.** Classification of a product under the Acceptance Program is subject to the conditions stated in the Agreement Governing Use of ADA Seal of Acceptance.
- **F.** Guidelines for the design and conduct of clinical studies are provided in Appendix I. Manufacturers interested in seeking the ADA Seal of Acceptance are encouraged to submit their clinical protocols to the Council for review prior to the start of clinical studies.

# 3. SUBMISSION MATERIALS

All submissions must include the following information based on product type and comply with the 'General Criteria for Acceptance' described in the Guidelines for Participation in the ADA Seal of Acceptance Program.

## A. Product Information

- i. Name of product(s)
- ii. Name of company



#### iii. FDA Documentation

- a) FDA registration and product listing must be provided.
- b) Evidence of FDA approval to market, if applicable (e.g., 510 (k) letter, pre-market approval, NDA/Evidence of FDA registration).

#### iv. Product Claims

a) Products approved under these category requirements will receive the following Seal bullet claim: helps clean removable prostheses. Data required to substantiate efficacy towards the Seal bullet claim is explained in Section C below. Please provide a list of all additional safety and efficacy claims beyond the Seal bullet claim. These claims should follow the ADA Advertising Standards and must undergo review and approval by the Council on Scientific Affairs before they can be included on product packaging. Substantiation for any health benefit claims, outside of the Seal bullet claims, must be provided through clinical and/or laboratory data specific to the product and is not addressed in Section C below. Whether clinical or laboratory data is required depends on the nature of the claim. For any questions regarding claim substantiation, please contact the ADA Seal Program.

# v. Product Specifications

- a) Chemical composition or components of the product and purpose of the various ingredients. To facilitate review, submitting the chemical composition, concentration, and purpose in tabular form is recommended.
- b) Material Safety Data Sheet (MSDS) (if applicable)
- c) Design of the product (if applicable)

## vi. Product Manufacturing

- Describe or list the quality procedures for manufacturing or testing of the product which demonstrate compliance with Good Manufacturing Practices.
- b) Certification of Good Manufacturing Practices can also be provided.

## vii. Product Instructions

- a) Include detailed instructions for product use.
- b) Include indications and contraindications for use, warnings, etc.

#### viii. Product Labeling/Packaging

a) All labeling/packaging should follow the ADA Advertising Standards and must be approved by the Council on Scientific Affairs before use. Companies may submit draft copy for approval. See iv. Product Claims above.



## ix. Product Samples

a) Submission requires three samples, one from three different production lots for analysis by the ADA Laboratories.

# **B.** Safety Data

- i. Evidence must be provided that the components of the product are safe for use in the oral cavity. When appropriate, standard toxicological, mutagenic, and/or carcinogenic testing may be required. Compliance with applicable FDA standards, as well as the Generally Recognized as Safe (GRAS) list should be provided (where appropriate).
- ii. For cleansers meant for use in the oral cavity, and in cases where new agents that do not appear on the GRAS list have been introduced into a cleanser, one six month clinical study may be required, which includes examinations of oral soft and hard tissues, toxicological studies, and microbiological profiles that should demonstrate that pathogenic or opportunistic microorganisms do not develop over the course of the study (See Appendix).
- iii. In products with agents with an established record of safe use in the oral cavity, clinical testing may not be required, but the manufacturer should provide supporting data and rationale for its use in the submitted product.
- iv. For cleansers meant for use outside of the oral cavity, laboratory testing data should be submitted showing that after the product has been rinsed off of the prosthesis following use, no dermal contact sensitization results due to residual product that might remain on the prosthesis. (See ANSI/ADA Standard No. 41 Evaluation of Biocompatibility of Medical Devices Used in Dentistry)
- v. Laboratory tests should be submitted to demonstrate that the cleanser will not adversely affect prostheses and their component materials under conditions of actual use (e.g. using ASTM D543-14 for plastics, and appropriate tests to show no etching, pitting or localized corrosion of metal components).

# C. Efficacy Data

- i. Supply one copy of all available physical and chemical property information developed in laboratory studies or similar materials that might be predictive of clinical use/behavior.
- ii. Two independent 30 day clinical studies should be submitted to support each of the efficacy claims for the product (e.g. removes stains; removes plaque/mouth film and soft debris) under conditions of actual use. Additional efficacy claims (e.g. antibacterial activity) must also be supported by appropriate studies. Additional guidelines are presented in the Appendix.
- **D. Supporting Literature:** Copies of the most significant articles or supporting literature demonstrating safety or efficacy of the product should be provided, where applicable.



# 4. REFERENCES

The following references were used in the development of these requirements and can be consulted for a more detailed discussion:

- ANSI/ADA Standard No. 41 Evaluation of Biocompatibility of Medical Devices Used in Dentistry, 2020
- ASTM D543-21 Standard Practices for Evaluating the Resistance of Plastics to Chemical Reagents
- ADA Advertising Standards: https://www.ada.org/publications/advertising-standards



# Appendix Clinical Protocol Guidelines for Cleansers for Removable Prostheses

The following guidelines are for the design and conduct of clinical studies for the evaluation of the safety and efficacy of denture cleanser products. Manufacturers are encouraged to submit their clinical protocols to the Council for review prior to the start of the clinical studies. The information indicated below is applicable to each independent clinical study.

## Sample Size

The Council recommends that a minimum of 30 subjects per test complete the study.

# **Subject Selection**

All subjects should be in good physical health with no medical problems that would contraindicate participation in the clinical study. Subjects should be screened for potential participation in the study and the screening pool should be examined for balance in terms of gender and broad age distribution. Subject population should be indicative of those for whom the product is intended. Subjects must refrain from the use of any non-study related denture cleansers. Other criteria for inclusion/exclusion of subjects must be provide.

# **Study Duration**

The Council recommends that two 30-day clinical studies are necessary to show safety and efficacy of the product. For products containing non-GRAS ingredients, a 6-month study for safety may be required.

#### Study Design

Each subject will have a complete oral cavity examination to determine eligibility for the study. The frequency of use of the product should be representative of actual use of the product in practice; and the user should be instructed in the proper use of the product, but not necessarily supervised. Measurements will be taken at baseline and at the conclusion of the study with an optional intermediate period. For those products with a directed use period, an observation immediately post-use is required.

#### **Safety Assessments**

Safety must be evaluated and acceptance is contingent upon demonstrating the absence of irreversible side effects resulting from the use of the product.

#### **Adverse Events**

All adverse events should be reported including altered oral sensations for each observation period e.g. burning mouth or altered taste).

#### Effect on oral tissues

Evidence that the product does not adversely affect oral soft issues should be provided. Subjects should be examined in the course of the study for the presence of pathologic conditions such as oral ulceration, candidiasis, or other secondary infections of the oral mucosa that may be manifestations of the proliferation of opportunistic microorganisms.

# **Efficacy Assessments**

Acceptance is contingent upon the results of laboratory tests used to validate each of the efficacy claims for the product under conditions of clinical use and behavior.

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