Shield frameworks; (6) misrepresenting whether CafePress would honor deletion requests; and (7) unfairly withholding commissions payable to shopkeepers.

The Proposed Orders contain provisions designed to prevent Respondents from engaging in the same or similar acts or practices in the future.

Summary of Proposed Order With Residual Pumpkin

Part I prohibits Residual Pumpkin from misrepresenting: (1) Privacy and security measures it takes to prevent unauthorized access to Personal Information; (2) the extent to which Residual Pumpkin is a member of any privacy or security program sponsored by a government, self-regulatory, or standard-setting organization; (3) privacy and security measures to honor users' privacy choices; (4) information deletion and retention practices; and (5) the extent to which it maintains and protects the privacy, security, availability, confidentiality, or integrity of Personal Information.

Part II requires Residual Pumpkin to establish and implement, and thereafter maintain, a comprehensive information security program ("Security Program") that protects the privacy, security, confidentiality, and integrity of Personal Information. Part III requires Residual Pumpkin to obtain initial and biennial data security assessments for 20 years. Part IV requires Residual Pumpkin to disclose all material facts to the assessor and prohibits Residual Pumpkin from misrepresenting any fact material to the assessment required by Part II. Part V requires Residual Pumpkin to submit an annual certification from a senior corporate manager (or senior officer responsible for its Security Program) that Residual Pumpkin has implemented the requirements of the order and is not aware of any material noncompliance that has not been corrected or disclosed to the Commission. Part VI requires Residual Pumpkin to notify the Commission of a "Covered Incident" within thirty days of discovering such incident.

Parts VII and VIII require Residual Pumpkin to pay to the Commission \$500,000 and describe the procedures and legal rights related to that payment. Part IX requires Residual Pumpkin to provide customer information to enable the Commission to administer consumer redress. Part X requires Residual Pumpkin to submit an acknowledgement of receipt of the order, including all officers or directors and employees having managerial responsibilities for conduct related to the subject matter of the order, and to

obtain acknowledgements from each individual or entity to which a Residual Pumpkin has delivered a copy of the order.

Part XI requires Residual Pumpkin to file compliance reports with the Commission and to notify the Commission of bankruptcy filings or changes in corporate structure that might affect compliance obligations. Part XII contains recordkeeping requirements for accounting records, personnel records, consumer correspondence, advertising and marketing materials, and claim substantiation, as well as all records necessary to demonstrate compliance with the order. Part XIII contains other requirements related to the Commission's monitoring of Respondent's order compliance.

Part XIV provides the effective dates of the order, including that, with exceptions, the order will terminate in twenty (20) years.

Summary of Proposed Order With PlanetArt

Part I prohibits PlanetArt from misrepresenting: (1) Privacy and security measures it takes to prevent unauthorized access to Personal Information; (2) the extent to which PlanetArt is a member of any privacy or security program sponsored by a government, self-regulatory, or standard-setting organization; (3) privacy and security measures to honor users' privacy choices; (4) information deletion and retention practices; and (5) the extent to which it maintains and protects the privacy, security, availability, confidentiality, or integrity of Personal Information.

Part II requires PlanetArt to establish and implement, and thereafter maintain, a comprehensive information security program that protects the privacy, security, confidentiality, and integrity of Personal Information. Part III requires PlanetArt to obtain initial and biennial data security assessments for 20 years. Part IV requires PlanetArt to disclose all material facts to the assessor and prohibits PlanetArt from misrepresenting any fact material to the assessment required by Part II.

Part V requires PlanetArt to submit an annual certification from a senior corporate manager (or senior officer responsible for its Security Program) that PlanetArt has implemented the requirements of the order and is not aware of any material noncompliance that has not been corrected or disclosed to the Commission. Part VI requires PlanetArt to notify the Commission of a "Covered Incident" within thirty days of discovering such incident. Parts VII

requires PlanetArt to provide notice to consumers to inform them of the breach and the settlement with the FTC.

Part VIII requires PlanetArt to submit an acknowledgement of receipt of the order, including all officers or directors and employees having managerial responsibilities for conduct related to the subject matter of the order, and to obtain acknowledgements from each individual or entity to which a PlanetArt has delivered a copy of the order.

Part IX requires PlanetArt to file compliance reports with the Commission and to notify the Commission of bankruptcy filings or changes in corporate structure that might affect compliance obligations. Part X contains recordkeeping requirements for accounting records, personnel records, consumer correspondence, advertising and marketing materials, and claim substantiation, as well as all records necessary to demonstrate compliance with the order. Part XI contains other requirements related to the Commission's monitoring of PlanetArt's order compliance.

Part XII provides the effective dates of the order, including that, with exceptions, the order will terminate in 20 years.

The purpose of this analysis is to facilitate public comment on the Proposed Orders, and it is not intended to constitute an official interpretation of the complaint or Proposed Orders, or to modify the Proposed Orders' terms in any way.

By direction of the Commission.

April J. Tabor,

Secretary.

[FR Doc. 2022–06022 Filed 3–21–22; 8:45 am]

BILLING CODE 6750–01–P

# GENERAL SERVICES ADMINISTRATION

[Notice MG-2022-01; Docket No. 2022-0002; Sequence No. 1]

Office of Federal High-Performance Green Buildings; Green Building Advisory Committee; Notification of Upcoming Web-Based Public Meeting

**AGENCY:** Office of Government-wide Policy, General Services Administration (GSA).

**ACTION:** Notice of public meeting.

**SUMMARY:** Notice of this web-based public meeting is being provided in accordance with the Federal Advisory Committee Act. This notice provides the date for the Green Building Advisory

Committee meeting, which is open to the public. Interested individuals must register to attend as instructed below under SUPPLEMENTARY INFORMATION.

**DATES:** The Green Building Advisory Committee will hold a web-based public meeting on Monday, April 18, 2022 from 11:00 a.m. to 5:00 p.m. Eastern Time (ET).

#### FOR FURTHER INFORMATION CONTACT: Dr.

Ken Sandler, Designated Federal Officer, Office of Federal High-Performance Green Buildings, Office of Government-wide Policy, General Services Administration, 1800 F Street NW (Mail-code: MG), Washington, DC 20405, at ken.sandler@gsa.gov or 202—219—1121. Additional information about the Committee, including meeting materials and agendas, will be available on-line at https://www.gsa.gov/gbac.

### SUPPLEMENTARY INFORMATION:

#### **Procedures for Attendance**

Contact Dr. Ken Sandler at ken.sandler@gsa.gov to register to attend this public web-based meeting. To register, submit your full name, organization, email address and phone number. Requests to attend the webbased meeting must be received by 5:00 p.m. ET, on Tuesday, April 12, 2022. Meeting call-in information will be provided to interested parties who register by the deadline. (GSA will be unable to provide technical assistance to any listener experiencing technical difficulties. Testing access to the webbased meeting site before the meetings is recommended.) Contact Dr. Sandler to register to provide public comment during the April 18, 2022 meeting public comment period. Attendees registered to provide public comment will be allowed a maximum of five minutes each and will need to provide written copies of their comments. Requests to provide public comment at the Committee meeting must be received by 5:00 p.m. ET, on Tuesday, April 12, 2022. To request for an accommodation, such as closed captioning, or to ask about accessibility, please contact Mr. Bryan Steverson at bryan.steverson@gsa.gov by Monday, April 4, 2022 to give GSA as much time as possible to process the request.

### Background

The Administrator of GSA established the Committee on June 20, 2011 (Federal Register/Vol. 76, No. 118) pursuant to Section 494 of the Energy Independence and Security Act of 2007 (42 U.S.C. 17123). Under this statute, the Committee provides independent policy advice and recommendations to GSA to advance federal building

innovations in planning, design, and operations to reduce costs, enable agency missions, enhance human health and performance, and minimize environmental impacts.

## April 18, 2022 Meeting Agenda

- Updates and Introductions
- Update on Embodied Carbon
- Environmental Justice and Equity for Federal Green Buildings Task Group: Proposed Advice Letter
- Federal Building Decarbonization Task Group: Proposed Advice Letter and Update
- Executive Order 14057: Update and Discussion
- New Committee Topics and Directions
- Public Comment
- Next Steps and Closing Comments

### Kevin Kampschroer,

Federal Director, Office of Federal High-Performance Green Buildings, Office of Government-wide Policy, General Services Administration.

[FR Doc. 2022–06040 Filed 3–21–22; 8:45 am]

BILLING CODE 6820-14-P

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Centers for Medicare & Medicaid Services

[CMS-3422-N]

Announcement of the Re-Approval of the American Association for Laboratory Accreditation (A2LA) as an Accreditation Organization Under the Clinical Laboratory Improvement Amendments of 1988

**AGENCY:** Centers for Medicare & Medicaid Services (CMS), Department of Health and Human Services (HHS). **ACTION:** Notice.

SUMMARY: This notice announces the application of the American Association for Laboratory Accreditation (A2LA) for approval as an accreditation organization for clinical laboratories under the Clinical Laboratory Improvement Amendments of 1988 (CLIA) program. We have determined that the A2LA meets or exceeds the applicable CLIA requirements. In this notice, we announce the approval and grant the A2LA deeming authority for a period of 6 years.

**DATES:** The approval announced in this notice is effective from March 23, 2022, until March 22, 2028.

**FOR FURTHER INFORMATION CONTACT:** Cindy Flacks, 410–786–6520.

SUPPLEMENTARY INFORMATION:

### I. Background and Legislative Authority

On October 31, 1988, the Congress enacted the Clinical Laboratory Improvement Amendments of 1988 (CLIA) (Pub. L. 100-578). CLIA amended section 353 of the Public Health Service Act. We issued a final rule implementing the accreditation provisions of CLIA on July 31, 1992 (57 FR 33992). Under those provisions, we may grant deeming authority to an accreditation organization if its requirements for laboratories accredited under its program are equal to or more stringent than the applicable CLIA program requirements in 42 CFR part 493 (Laboratory Requirements). Subpart E of part 493 (Accreditation by a Private, Nonprofit Accreditation Organization or Exemption Under an Approved State Laboratory Program) specifies the requirements an accreditation organization must meet to be approved by CMS as an accreditation organization under CLIA.

# II. Notice of Approval of A2LA as an Accreditation Organization

In this notice, we approve the American Association for Laboratory Accreditation (A2LA) as an organization that may accredit laboratories for purposes of establishing their compliance with CLIA requirements in all specialties and subspecialties. We have examined the initial A2LA application and all subsequent submissions to determine its accreditation program's equivalency with the requirements for approval of an accreditation organization under subpart E of part 493. We have determined that the A2LA meets or exceeds the applicable CLIA requirements. We have also determined that the A2LA will ensure that its accredited laboratories will meet or exceed the applicable requirements in subparts H, I, J, K, M, Q, and the applicable sections of subpart R of part 493. Therefore, we grant the A2LA approval as an accreditation organization under subpart E of part 493, for the period stated in the DATES section of this notice for all specialties and subspecialties under CLIA. As a result of this determination, any laboratory that is accredited by the A2LA during the time period stated in the DATES section of this notice will be deemed to meet the CLIA requirements for the listed specialties and subspecialties, and therefore, will generally not be subject to routine inspections by a State survey agency to determine its compliance with CLIA requirements. The accredited laboratory,