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Date: April 18, 2023 Information Only

To: CSU Procurement Offices

CSU Risk Management

**CSU Sponsored Programs Administration** 

**CSU Research Officers** 

From: David Beaver, Chief Procurement Officer David Beaver

**Subject:** Guidance on Purchasing Gene Synthesis Equipment or Products

### **Purpose**

This Guidance Memo informs the research community about Assembly Bill (AB) 1963 and provides guidance on purchasing gene synthesis equipment or gene synthesis products from gene synthesis providers.

University of California Office of the President developed AB 1963 guidance in coordination with the California State University (CSU) Office of the Chancellor. Representatives from Procurement, Environmental Health and Safety, Sponsored Programs Administration and the Office of Research participated in the development and review process.

### **Background**

Gene synthesis is the process of designing and synthesizing sequences of nucleic acids, which allows individuals to create a gene from scratch. Synthetic genes are used for research to test genetic hypotheses, create advanced gene-editing systems, identify new drugs, and even develop life-saving vaccines. While gene synthesis is vital for research, there are biosecurity concerns associated with unauthorized access to potentially harmful biological agents.

The U.S. has made progress to reduce biosecurity risks associated with individuals with ill intent gaining access to harmful biological agents. In 2010, the US Department of Health and Human Services (HHS) published <u>guidance for commercial gene synthesis providers</u> that recommended screening sequence orders and customers. In addition, the industry-led <u>International Gene Synthesis Consortium</u> (IGSC) was formed in 2009 to share sequence and customer screening methods among its members. IGSC members commit to following



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specified gene sequence and customer screening protocols as well as record keeping and reporting requirements (see IGSC's <u>Harmonized Screening Protocol</u>, most recently updated in 2017). The federal government continues to evaluate biosafety concerns and provide updated guidance.

In August 2022, California passed, and the Governor signed <u>AB 1963</u>, which amends sections 66360 and 66361 of the Education Code to require the CSU and to request that UC develop systemwide guidance for purchasing gene synthesis equipment or gene synthesis products from providers who prevent the misuse of synthetic genes and safeguard the benefits of gene synthesis technology while minimizing risk.

#### Guidance

Pursuant to AB 1963, UC and CSU are jointly issuing guidance to their respective University campus communities who may be involved in purchasing gene synthesis equipment or gene synthesis products.

Prior to purchasing gene synthesis equipment or gene synthesis products, those placing orders, including researchers, lab managers, students, and local procurement offices, should ensure that the gene synthesis provider meets the IGSC criteria or that the provider applies protections commensurate to the IGSC criteria. An explanation of the IGSC criteria and commensurate protections are provided in the subsequent sections in this document.

Gene synthesis providers may complete an attestation form to demonstrate that they apply the IGSC criteria or commensurate protections. Alternatively, gene synthesis providers may present other written information demonstrating that they prevent the misuse of synthetic genes and safeguard the benefits of gene synthesis technology while minimizing risk. UC Systemwide Procurement has created a Smartsheet dashboard listing which gene synthesis providers have met the IGSC criteria. This Smartsheet dashboard includes an online tool for requesting provider attestations and is available for use by both UC and CSU. In addition, a sample paper attestation form is provided in Appendix I of this guidance document.

CSU purchasing offices should keep gene synthesis providers' attestation letters or other written information on file within purchasing systems so that a gene synthesis provider will only have to provide this information once. For this reason, the preferred repository for requesting, receiving, and verifying supplier attestations is the UC Smartsheet tool. Those purchasing synthesis equipment or gene synthesis products should place orders from gene synthesis providers who have a complete online attestation, attestation letter or other written information on file. Additionally, suppliers who are members of the <a href="International Gene Synthesis">International Gene Synthesis</a> Consortium (IGSC) shall be considered as "following ISGC protocol" regardless of attestation status.



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#### The IGSC Criteria

The IGSC is an industry-led group of gene synthesis companies and organizations formed to design and apply a common protocol to screen both the sequences of synthetic gene orders and the customers who place them.

The IGSC's "<u>Harmonized Screening Protocol for Gene Sequence & Customer Screening to Promote Biosecurity</u>" establishes five core components that IGSC companies must apply to promote the safe use of synthetic genes:

- Gene Sequence Screening: The complete DNA sequence of every synthetic gene order is to be screened against a Regulated Pathogen Database developed by the consortium and one or more of the internationally coordinated sequence reference databanks (i.e., NCBI/GenBank, EBI/EMBL or DDBJ). Amino acid sequences of possible translation products for each synthetic gene ordered will also be screened.
- Gene Customer Screening: A complete and thorough screening of each potential gene synthesis customer will be conducted to establish identity and clearance for delivery of genes ordered, in accordance with national guidelines. The screening protocol assigns special considerations to the ordering of Select Agent genes.
- Record Keeping: The IGSC companies will keep all screening, customer and order records for at least eight years.
- Order Refusal & Reporting: The IGSC companies reserve the right to refuse to fill any
  order and to notify authorities upon identifying potentially problematic orders,
  coordinating efforts with local and national law enforcement and intelligence agencies.
- Regulatory Compliance: The IGSC companies comply with all applicable laws and regulations governing the synthesis, possession, transport, export and import of gene synthesis and other products.

#### **Commensurate Protections**

Gene synthesis providers who do not apply the IGSC criteria may attest to meeting protections commensurate to the IGSC specific to gene sequencing screening and gene customer screening.

In terms of gene sequencing screening, providers should have a way to screen requested sequences to identify if a request is a "sequence of concern."



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The 2010 HHS Screening Framework Guidance for Providers of Synthetic Double-Stranded DNA<sup>1</sup> provides a set of recommended practices to companies synthesizing double-stranded (ds) DNA to encourage such companies to screen both their customers and requested sequences. HHS explains that "[t]he purpose of sequence screening is to identify when "sequences of concern" are ordered. Identification of a "sequence of concern" does not necessarily imply that the order itself is of concern. Rather, when a "sequence of concern" is ordered, further follow-up procedures should be used to determine if filling the order would raise concern. Sequence screening is recommended for all dsDNA orders."

HHS defines "sequences of concern" as sequences derived from or encoding select agents and toxins or items on the CCL, except when also found in unregulated organisms; or sequences that contribute to toxicity or pathogenicity, whether derived from or encoding regulated or unregulated biological agents.

As to gene customer screening, HHS explains that the purpose of customer screening is to verify the legitimacy of the customer and the principal user, to confirm that the customer and principal user placing an order are acting within their authority, and to verify the legitimacy of the end-use.

### Resources for Compliance with AB 1963

As a resource for the UC and CSU community, UC Systemwide Procurement developed a Smartsheet dashboard listing which gene synthesis providers have met the IGSC criteria or attested to providing protections commensurate to the IGSC criteria. This tool may also be used to send requests to gene synthesis providers to complete an attest of either meeting the ICSC criteria or providing commensurate protections. Prior to purchasing gene synthesis equipment or gene synthesis products, those placing orders for these equipment or products are encouraged to use this tool to either check for compliance with AB 1963 or requesting attestation forms.

The Smartsheet dashboard and tool for verifying and requesting provider attestations may be accessed here.

As an additional resource, a sample paper attestation form is provided in Appendix I of this guidance document. The attestation form may be used as a reference. Gene synthesis providers may also use this form to complete the attestation; however, UC and CSU recommend completing the attestation via the web tool.

<sup>&</sup>lt;sup>1</sup> In 2020 and 2022, HHS issued requests for information to update the 2010 Screening Framework Guidance for Providers of Synthetic Double-Stranded DNA. To date of this RPAC Guidance Memo, the HHS Screening Framework has not been updated. More information on these efforts can be found <a href="https://example.com/here-repeated-new-repeated-

### **Additional General Information on Biosafety**

While AB 1963 facilitates measures concerning the procurement of gene synthesis equipment or gene synthesis products, the research community is reminded that they must follow all applicable laws and policies regarding biosafety, including submitting a Biological Use Authorization if applicable, if they are working with certain kinds of biological agents, such as recombinant DNA, synthetic nucleic acids, and pathogens.

Campuses Institutional Biosafety Committees (IBC) are responsible for enforcing policies and guidelines related to university-related use of all potentially hazardous biological agents, including but not limited to infectious agents, human and non-human primate materials (including established cell lines), CDC select agents, recombinant DNA and studies involving human gene transfer. The Committee ensures that research involving these agents is conducted in a manner that does not endanger the researcher, laboratory worker, human research subjects, the public or the environment.

The National Institutes of Health (NIH has in place <u>Guidelines for Research Involving</u> <u>Recombinant or Synthetic Nucleic Acid Molecules</u> that applies to researchers who receive NIH funding. The NIH Guidelines provides details on safety practices and containment procedures for basic and clinical research involving recombinant or synthetic nucleic acid molecules, including the creation and use of organisms and viruses containing recombinant or synthetic nucleic acid molecules.

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# Appendix I

## **Sample Attestation Form**

In August 2022, California passed <u>AB 1963</u> , which amends sections 66360 and 66361 of the Education Code to require CSU and to request that UC develop systemwide guidance for purchasing gene synthesis equipment or gene synthesis products from providers who prevent the misuse of synthetic genes and safeguard the benefits of gene synthesis technology while minimizing risk.
My name is on behalf of company I understand that the University is relying on this attesting prior to purchasing gene synthesis equipment or gene synthesis products.
[Check One Below]  ☐ My company follows the International Gene Synthesis Consortium Criteria for Harmonized Screening Protocol.
□ My company carries out gene customer screenings and gene sequencing screenings to identify when "sequences of concern" are ordered. Identification of a "sequence of concern" does not necessarily imply that the order itself is of concern. Rather, when a "sequence of concern" is ordered, further follow-up procedures are taken to determine if filling the order would raise concern.
Should the information checked above change, providers are expected to give updated information to the University.
Company
Printed name
Title
Signed
Date