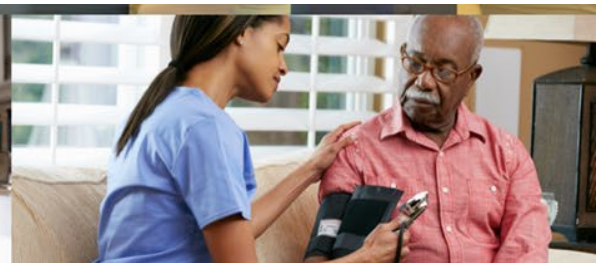




Pathways to Health Equity

Grantee Technical Assistance Session #3

June 1, 2022



Objectives for Today's Session

- CHRC expectations for data collection
 - Unique patient identifier and demographic information to be collected
 - Definition of patients served / unduplicated patients served
- Overview of CRISP panel uploading
- Topics of future technical assistance sessions- open discussion
- Questions & Answers period

Unique Patient Identifier & Demographic Information to be Collected

Pathways grantees will be expected to collect the following information as part of their Patient Intake form.

- Unique Participant Identifier (Patient_ID, MRN, etc.)
- First Name
- Last Name
- DOB
- Address
- Gender
- Race
- Ethnicity

If you already collect this information and have developed your own patient intake form, please forward a copy to CHRC staff for final approval.

Definition of Patient Served

The definition document will be distributed and discussed with Pathways grantees.

We recognize that this definition may impact information contained in your milestones and deliverables report.

CHRC and CRISP staff will work with Pathways grantees to finalize your milestones and deliverables report.

Topics of Future TA Sessions- open discussion

- **June 15** – CHRC Grant Monitoring, Consumer Engagement / Community Outreach, Marketing, Implementation
- **June 29** – CHRC Administrative topics – grant modifications, subcontracts/MOUs; fiscal documentation
- **July 13** – CRISP SDOH tools
- **Additional TA Sessions to be scheduled**

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