

BAME Commission

Project plan: research on causes of health inequalities amongst BAME communities

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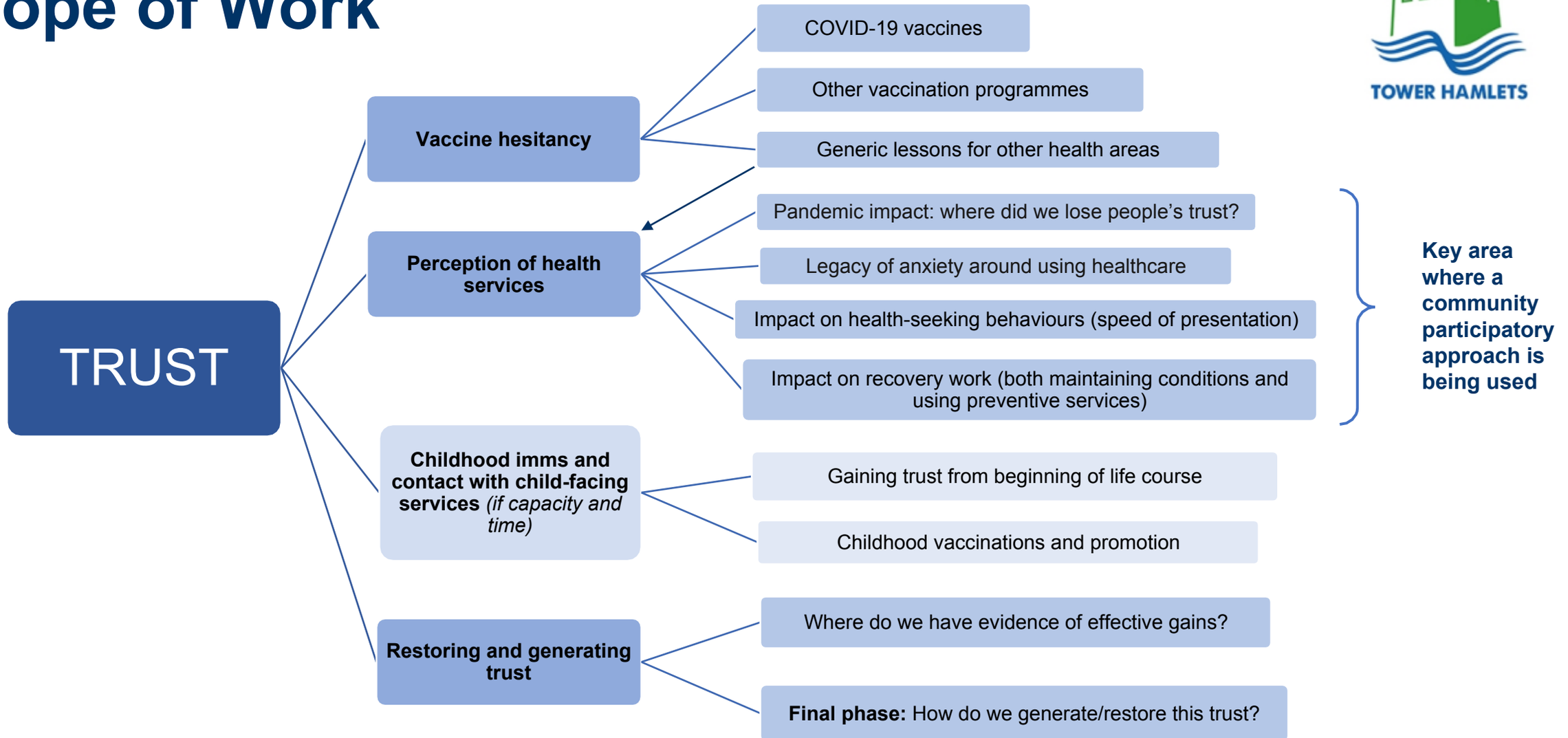
Recommendation:



“The Health and Wellbeing Board, by the end of 2021, undertake detailed external research on causes of health inequalities amongst BAME communities which puts engagement of the community at the forefront of its work to identify issues and solutions”



Scope of Work



DISAGGREGATION OF DATA INTO GRANULAR ETHNIC GROUPS



Research Phases



Phase 1: *“What we already have”*

- Synthesise information already collected from relevant literature
- Analyse qualitative insights data (collected by Public Health team) by project scope topics
- Integrate findings from existing vaccine hesitancy work (i.e. West Co, Social Action for Health, Community Navigators etc.)

Phase 2: *“Additional data to collect”*

- Health data to complement the community participatory work around perceptions of services
- Focused pieces of additional work which need to be done around vaccine hesitancy
- Service provider engagement

Phase 3: *“Community-based participatory research”*

- Generating additional insight; this will then be built on by co-developing focused actions
- Primary focus a) carrying out a deep dive in to the perception of health services and how the experience throughout Covid has impacted on trust b) to consider how grief and loss has impacted on the relationship of trust, and how we can begin to build up trust through this void of grief and trauma c) to explore how residents feel that their ethnicity impacts on how they use and perceive health services and d) to generate actions on how trust can be restored between BAME residents and services

