The HiLo Trial

Myles Wolf, MD, MMSc
The HiLo Trial
Myles Wolf, MD, MMSc
What is a Pragmatic Trial?

“The overall goal of the Collaboratory is to strengthen the national capacity to implement cost-effective large-scale research studies that engage health care delivery organizations as research partners.”
What is a Pragmatic Trial?

- “Real world” trial: evaluates effectiveness rather than efficacy

“The overall goal of the Collaboratory is to strengthen the national capacity to implement cost-effective large-scale research studies that engage health care delivery organizations as research partners.”
What is a Pragmatic Trial?

- “Real world” trial: evaluates effectiveness rather than efficacy
- Non-restrictive eligibility criteria

“The overall goal of the Collaboratory is to strengthen the national capacity to implement cost-effective large-scale research studies that engage health care delivery organizations as research partners.”
What is a Pragmatic Trial?

- “Real world” trial: evaluates effectiveness rather than efficacy
- Non-restrictive eligibility criteria
- Embedded in clinical care delivery

“The overall goal of the Collaboratory is to strengthen the national capacity to implement cost-effective large-scale research studies that engage health care delivery organizations as research partners.”
What is a Pragmatic Trial?

• “Real world” trial: evaluates effectiveness rather than efficacy
• Non-restrictive eligibility criteria
• Embedded in clinical care delivery
• Intervention implemented by clinical care providers

“The overall goal of the Collaboratory is to strengthen the national capacity to implement cost-effective large-scale research studies that engage health care delivery organizations as research partners.”
What is a Pragmatic Trial?

- “Real world” trial: evaluates effectiveness rather than efficacy
- Non-restrictive eligibility criteria
- Embedded in clinical care delivery
- Intervention implemented by clinical care providers
- Outcomes ascertained using clinically acquired data

“The overall goal of the Collaboratory is to strengthen the national capacity to implement cost-effective large-scale research studies that engage health care delivery organizations as research partners.”
What is a Pragmatic Trial?

- “Real world” trial: evaluates effectiveness rather than efficacy
- Non-restrictive eligibility criteria
- Embedded in clinical care delivery
- Intervention implemented by clinical care providers
- Outcomes ascertained using clinically acquired data
- Noise is expected

“The overall goal of the Collaboratory is to strengthen the national capacity to implement cost-effective large-scale research studies that engage health care delivery organizations as research partners.”
What is a Pragmatic Trial?

- “Real world” trial: evaluates effectiveness rather than efficacy
- Non-restrictive eligibility criteria
- Embedded in clinical care delivery
- Intervention implemented by clinical care providers
- Outcomes ascertained using clinically acquired data
- Noise is expected

Result: **generalizable findings, sustainable intervention, efficient conduct**

“The overall goal of the Collaboratory is to strengthen the national capacity to implement cost-effective large-scale research studies that engage health care delivery organizations as research partners.”
## Collaboratory trials

<table>
<thead>
<tr>
<th>Round 1</th>
<th>Round 2</th>
<th>Round 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Suicide Prevention using Electronic Engagement</td>
<td>Increasing Advance Care Directives in Nursing Homes using Videos</td>
<td>Increasing Advance Care Planning in Oncology with Videos and Clinician Training</td>
</tr>
<tr>
<td>Increasing Colorectal Screening with Automated Reminders</td>
<td>Increasing Implementation of Practice Guidelines for DM, HTN, CKD Using Practice Facilitators</td>
<td>Improving Adolescent Behavior with Parent Training</td>
</tr>
<tr>
<td>Multidisciplinary Management of Opioid Dependence</td>
<td>Reducing PTSD in Trauma Victims Using ER Intervention</td>
<td>Increasing ER Initiation of Buprenorphine for Opioid Use Disorder</td>
</tr>
<tr>
<td>Prevent MRSA Bacteremia</td>
<td></td>
<td>Improving Adherence to CV Medications Using Electronic Nudges</td>
</tr>
<tr>
<td>Reducing Unnecessary Imaging Studies by Adding Epi Info to Reports</td>
<td></td>
<td>Increasing Palliative Care in the ER HiLo</td>
</tr>
<tr>
<td>TiME: Reducing Mortality in ESRD with Longer HD</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Collaboratory trials

**Round 1**
- Suicide Prevention using Electronic Engagement
- Increasing Colorectal Screening with Automated Reminders
- Multidisciplinary Management of Opioid Dependence
- Prevent MRSA Bacteremia
- Reducing Unnecessary Imaging Studies by Adding Epi Info to Reports
- TiME: Reducing Mortality in ESRD with Longer HD

**Round 2**
- Increasing Advance Care Directives in Nursing Homes using Videos
- Increasing Implementation of Practice Guidelines for DM, HTN, CKD Using Practice Facilitators
- Reducing PTSD in Trauma Victims Using ER Intervention

**Round 3**
- Increasing Advance Care Planning in Oncology with Videos and Clinician Training
- Improving Adolescent Behavior with Parent Training
- Increasing ER Initiation of Buprenorphine for Opioid Use Disorder
- Improving Adherence to CV Medications Using Electronic Nudges
- Increasing Palliative Care in the ER HiLo
ESRD: Trials needed & an ideal setting for pragmatism

- ESRD population is desperately in need of clinical innovation
  - High event rates
  - Few, if any therapies proven by RCT
- Highly accessible study population with 3 x weekly clinical encounters
- Highly granular, regular, uniform data collected in routine clinical care → EHR
  - Remote biochemical monitoring
  - Pragmatic ascertainment of outcomes, covariates
- Centralized infrastructure of dialysis provider organizations allows for
  - Centralized implementation
  - Inclusion of large number of facilities with broad geographic distribution
  - Facility-level randomization
Scientific basis of HiLo

- End-stage renal disease (ESRD): ~500,000 in US alone
- Unacceptably high rates of:
  - Hospitalization: ~2 per patient-year
  - Mortality: 15–20%
  - Driven primarily by CVD, but CVD interventions failed in ESRD trials
- Focus on ESRD-specific risk factors: Hyperphosphatemia
  - Ubiquitous complication of ESRD
  - Experimental data: causal effects on arterial calcification, LVH, high PTH, FGF23
  - Hyperphosphatemia & all of above associated with CVD, death
- Based on preclinical & observational human data, opinion-based guidelines suggest P <5.0 mg/dl using P binders, low P diet

Block GA, AJKD 1998; Block GA, JASN 2004
Clinical equipoise

• Lack of RCTs
  • Target phosphate unproven by RCT
  • No RCT tested the effects of FDA-approved phosphate binders on clinical outcomes
• Major unanswered questions:
  1. Do phosphate binders, as currently deployed, improve outcomes in ESRD?
  2. Does lowering serum P towards normal improve outcomes in ESRD?
• Hidden risks – excessive treatment to unnecessarily low target may worsen outcomes:
  • Paradoxically increasing risk by inducing calcium, lanthanum or iron overload
  • Causing GI side effects that exacerbate malnutrition
  • Eroding patients’ QOL: adding P-related demands to high pill burdens
• Potential risks may have escaped detection precisely because of lack of RCTs
Hypotheses

1. **Primary**: Compared to the current standard approach of targeting serum phosphate levels of <5.5 mg/dl, less stringent control of serum phosphate to target levels of 6–7 mg/dl will yield non-inferior rates of all-cause hospitalization among patients with ESRD undergoing hemodialysis.

2. **Main secondary**: Compared to strict phosphate control, less stringent control will reduce risk of all-cause mortality, enhance markers of diet and nutrition, and improve quality of life.
Overview of HiLo

Pragmatic, multicenter, cluster-randomized, open-label, non-inferiority, outcomes trial

• Compare effects of two different phosphate management strategies
  • Liberal P control, targeting 6–7 mg/dl, or
  • Strict P control, targeting <5.5 mg/dl
  • Facility-level cluster randomization: simplify trial execution, prevent within-facility “bleeding” of intervention arms, support remote study monitoring

• N = ~4400 patients being treated with hemodialysis at >100 facilities

• Partners
  1. Large national for-profit dialysis corporation: DaVita, Inc.
  3. Small regional academic program: University of Utah

• Build on lessons learned from the TIME trial
Outcomes

1. Primary
   • All-cause hospitalization rate: total counts per person-years of follow-up (continuous)

2. Main secondary
   • All-cause mortality, time-to-event (superiority)

3. Other
   • Total inpatient hospital days per person-years of follow-up
   • Cause-specific hospitalizations in Medicare beneficiaries based on merging clinical data from HiLo with claims data from the CMS Virtual Research Data Center as in PROVEN
   • Diet & nutrition: serum albumin, protein catabolic rate (PCR)
   • Quality of life: 36-SF
Justification of all-cause hospitalization as primary outcome

- Important to all ESRD stakeholders: patients, families, clinicians, dialysis providers, payers
- For many patients, avoiding hospitalization/enhance QOL >>> prolong survival
- HyperP contributes to multiple complications → hospitalization
- Somewhat “noisy”:
  - Includes events unlikely to be influenced by P control
  - But... more frequent hospitalization: marker of poor health, correlates with mortality in ESRD
  - All-cause mortality: subject to the same noisiness.
  - To address noise, HiLo powered to detect a small effect size
- Hospitalization: accepted endpoint in other areas
- Dialysis providers: near 100% complete data about dates/duration of hospitalizations; not causes
- Collecting real-time data on all-cause hospitalization eliminates events adjudication
- Continuous variable: more desirable statistically than time-to-event outcomes
Eligibility Criteria

Dialysis Facility:
• Willingness of MD director, nephrologists, dietitians to adopt either P target
• Willingness of facility managers to allow dietitian participation
• Facility dietitian willingness to implement trial, attend training teleconferences

Individual Participant Criteria:
• Liberal to simplify study, facilitate enrollment, maximize generalizability
• Adults >18 years of age treated with standard in-center maintenance HD
• Willing/able to provide written informed consent
• All vintages; pre-specified secondary analysis stratified by vintage < versus >3 years
Intervention arms: Phosphate titration

HiLo will develop two phosphate titration protocols with same “look and feel” as in practice

1. **Low serum phosphate target**
   - Current standard of care
   - Goal: titrate serum phosphate <5.5 mg/dl
   - Anticipate 4.8–5.2 mg/dl

2. **Higher serum phosphate**
   - Intervention strategy
   - Goal: titrate serum phosphate to 6–7 mg/dl
   - Anticipate 6.5–6.8 mg/dl
   - Since serum P = 4–7 mg/dl in most patients with ESRD, ≥1 mg/dl difference = ≥33% difference within modifiable range of time-averaged P exposure
   - Specific binder choices: discretion of local providers based on local practice
Preliminary pilot trials

Do we need informed consent?

Regulatory criteria for waiving consent

1. The research involves no more than minimal risk to the subjects
2. The waiver or alteration will not adversely affect the rights and welfare of the subjects
3. The research could not practicably be carried out without the waiver or alteration
4. Whenever appropriate, the subjects will be provided with additional pertinent information after participation

45 CFR Part 46 ("The Common Rule")
Do we need informed consent?

Regulatory criteria for waiving consent

1. The research involves no more than minimal risk to the subjects
2. The waiver or alteration will not adversely affect the rights and welfare of the subjects
3. The research could not practicably be carried out without the waiver or alteration
4. Whenever appropriate, the subjects will be provided with additional pertinent information after participation

45 CFR Part 46 ("The Common Rule")
Obtaining consent without on-site study staff

1. Paper consent
2. Electronic consent via internet-linked tablets
3. Centralized DCC-based phone resource

Benefits of Obtaining Consent
- Promote adherence
- Direct-push study updates, newsletter to participants
- Obtain additional data without CRFs or on-site study staff: e.g., PROs
Randomization

- Cluster-randomization of dialysis facilities using stratification to achieve balance across the two arms
- For the larger providers, we will stratify by:
  - Chain (DaVita or DCI)
  - Facility size (above or below the provider’s median facility census)
  - Tertiles of facility-specific standardized mortality rates (SMR): indicator of facilities’ global severity of illness and clinical outcomes
Overview of HiLo informatics
Data Collection

1. Demographic and comorbidity data at study entry:
   - Dialysis facility zip code
   - Age
   - Sex
   - Race
   - Ethnicity
   - Height
   - Weight
   - Dialysis vintage
   - Co-morbid illnesses noted on admission to the dialysis facility (ICD-9/10 codes)
   - Cause of end-stage renal disease
Data Collection

2. Dialysis treatment data:
   - Dialysis adequacy, Kt/V: once per month
   - Vascular access type (presence or absence of catheter): once per month

3. Health-Related Quality of Life
   - Quality of life survey
   - PROs: TBD
4. Laboratory Data: monthly
   - Hemoglobin
   - Albumin
   - Calcium
   - Phosphate
   - Protein catabolic rate
   - Serum ferritin
   - Transferrin saturation

5. Laboratory Data: once every 3 months
   - Intact parathyroid hormone
Data Collection

7. Hospitalizations data: all
8. Medications: all
   - Phosphate binders
   - Activated vitamin D
   - Home medications
9. Status Change: all
   - Date of transfer to another dialysis facility
   - Date of kidney transplantation
   - Date of transfer to peritoneal dialysis
   - Date of withdrawal from dialysis
   - Date of death
Sample size, primary analysis

• **Primary analysis**: comparison of per-person year (PPY) hospitalization rates between P target arms
  - Null hypothesis: low P target is superior (lower hospitalization rate)
  - Alternative hypothesis: high P target is non-inferior

• **Sample size assumptions**
  - 45-month study, 18-month enrollment period
  - 2.0 PPY mean hospitalization rate
  - Non-inferiority delta of 0.114 = 5% of 2.0 PPY mean hospitalization rate
  - SD of the hospitalization rate of 1.0, 1.2, or 1.4 for both groups
  - Annual loss to follow-up: 5%
  - Two-sided α: 0.05; Power: 80% or 90%
  - Mean enrollment of 30–35 patients per facility with SD of 0 and 9.5
  - Intra-class correlation coefficient between two patients from the same facility of 0.003

• Randomizing **120–150 facilities** (65–75/P target arm), **4400 patients** will provide >80% power to detect non-inferiority of the high P target
Sample size, secondary analysis

- **Main secondary analysis**: comparison of mortality rates between P target arms
  - Null hypothesis: survival curves are the same
  - Alternative hypothesis: higher mortality in one arm or the other
- **Sample size assumptions**
  - 45-month study; 18-month enrollment period
  - Annual mortality rate of 15%
  - Hazards ratio of mortality comparing P target arms: 0.8 and 0.85
  - Annual loss to follow-up: 5%
  - Two-sided $\alpha$: 0.05; Power: 80% or 90%
  - Mean enrollment of 30–35 patients per facility with SD of 0 and 9.5
  - Intra-class correlation coefficient between two patients from the same facility of 0.0012
- Randomizing **120–150 facilities** (65–75/P target arm), **4400 patients**, will provide >80% power to detect a HR of 0.85 for all-cause death between P target arms, >90% power to detect a HR of 0.80.
- Change from baseline in serum albumin, PCR, QOL, PROs: continuous variables; ample power
Safety

- Independent DSMB to be convened by NIDDK
- Given pragmatic design, we will not collect information on AEs as in traditional RCTs
- Will monitor relevant, already collected laboratory parameters:
  - Monthly serum phosphate, calcium, PTH.
- Will monitor for primary, secondary outcomes:
  - Will capture clinically important AEs through all-cause hospitalizations, all-cause death
- Since individual patients’ medical care is ultimate responsibility of primary providers, they may, at their discretion, reduce or temporarily discontinue P binders as in non-RCT setting, e.g.:
  - Hypercalcemia
  - GI symptoms
  - Hypophosphatemia
  - Participant preference
Data and Safety Monitoring Board (DSMB)

- NIDDK will convene a Data and Safety Monitoring Board (DSMB)
- DSMB will review reports containing data on:
  - Trial progress (enrollment, demographic, etc.)
  - Data quality
  - Rates of adverse events by masked treatment arm
  - Other safety data
  - Blinded primary and secondary efficacy outcomes (option to un-blinding)
- DSMB will make recommendations to NIDDK about progress, safety, and continuation
- Institutional Review Board (IRB) will also receive DSMB reports
- All sites will receive continuation letters from DSMB
Pivotal role of dietitians

• Dietitians: on-the-ground personnel who implement HiLo
  • Employed by dialysis organizations
  • Present in all dialysis units
  • Among the most motivated caregivers on dialysis teams
  • See all patients at least monthly
  • Serve as primary decision makers for titration of P-related Rx
  • Existing rapport with patients will facilitate adherence
  • Relying on clinical personnel to implement trial: consistent with pragmatism

• To engage dietitians in HiLo:
  • Recruit dietitian representatives to Steering Committee: input into design & implementation
  • Identify regional dietitian champions for HiLo
  • Drive interest via presentations at national meetings: nephrology, dialysis company, dietitians
Pragmatic features of HiLo

• Cluster randomization: enhance fidelity of interventions, simplify logistics
• Liberal eligibility criteria
• Internet/tablet-based eConsent for individual patient-level informed consent
• In-center dietitians implement the intervention
• Phosphate management protocols with same “look and feel” as in clinical practice
• Implementation of intervention using approved medications
• Use of EHR data to remotely & continuously monitor fidelity of interventions
• Use of EHRs to extract clinical data, outcomes
• Merge with Medicare claims for 2’ analyses
Possible HiLo results: Would rapidly influence ESRD practice

1. **Higher P target non-inferior:**
   - Contradicts guidelines
   - Relax P target, dietary restrictions
   - Reallocate dialysis resources
   - Reduce burden on patients

2. **Higher P target superior:**
   - Contradicts guidelines
   - Relax P target, dietary restrictions

3. **Higher P target inferior = low P target superior:**
   - Fail to reject null hypothesis
   - First definitive clinical trial-grade evidence for opinion-based guidelines for P management
   - For CMS: justify P as a validated dialysis quality-of-care measure
   - Support additional trials of P control in earlier stages of CKD
<table>
<thead>
<tr>
<th>Name</th>
<th>Affiliation</th>
<th>Role</th>
</tr>
</thead>
<tbody>
<tr>
<td>Myles Wolf, MD, MMSc</td>
<td>Duke University, Duke Clinical Research Institute</td>
<td>Principal Investigator, Steering Committee Chair</td>
</tr>
<tr>
<td>Srinivasan Beddu, MD</td>
<td>University of Utah Health</td>
<td>Steering Committee, Utah liaison</td>
</tr>
<tr>
<td>Geoffrey Block, MD</td>
<td>Denver Nephrology Associates</td>
<td>Steering Committee</td>
</tr>
<tr>
<td>Steven Brunelli, MD, MSCE</td>
<td>DaVita Clinical Research</td>
<td>Steering Committee, DaVita liaison</td>
</tr>
<tr>
<td>Hrishikesh Chakraborty, DrPH</td>
<td>Duke Clinical Research Institute</td>
<td>Steering Committee, Statistical lead</td>
</tr>
<tr>
<td>Laura Dember, MD</td>
<td>University of Pennsylvania</td>
<td>Steering Committee</td>
</tr>
<tr>
<td>Tamara Isakova, MD, MMSc</td>
<td>Northwestern University</td>
<td>Steering Committee</td>
</tr>
<tr>
<td>Matthew Roe, MD</td>
<td>Duke Clinical Research Institute</td>
<td>Steering Committee</td>
</tr>
<tr>
<td>Daniel Weiner, MD, MS</td>
<td>Dialysis Clinic, Inc.</td>
<td>Steering Committee, Dialysis Clinic, Inc. liaison</td>
</tr>
<tr>
<td>Becky Brosch, RD, CSR, LD</td>
<td>DaVita</td>
<td>Steering Committee, Dietitian Co-lead</td>
</tr>
<tr>
<td>Dietitian, TBD</td>
<td>Dialysis Clinic, Inc.</td>
<td>Steering Committee, Dietitian Co-lead</td>
</tr>
<tr>
<td>Dietitian, TBD</td>
<td>University of Utah Health</td>
<td>Steering Committee, Dietitian Co-lead</td>
</tr>
<tr>
<td>Laura Johnson, Davy Andersen</td>
<td>Duke Clinical Research Institute</td>
<td>Project Leader</td>
</tr>
<tr>
<td>Andrew Mackelfresh</td>
<td>Duke Clinical Research Institute</td>
<td>Bioinformatics lead</td>
</tr>
</tbody>
</table>
Questions & Comments