**Additional file 1: Table S1**: The World Health Organization (WHO) Trial Registration Data Set for the SIMPLER study.

<table>
<thead>
<tr>
<th>Data category</th>
<th>Information</th>
</tr>
</thead>
<tbody>
<tr>
<td>Countries of Recruitment</td>
<td>Australia</td>
</tr>
<tr>
<td>--------------------------</td>
<td>-----------</td>
</tr>
<tr>
<td>Health Condition(s) or Problem(s) Studied</td>
<td>Polypharmacy, Medication incidents, Dementia, Cognitive impairment, Falls, Hospitalisation, Frailty</td>
</tr>
<tr>
<td>Intervention(s)</td>
<td>Medications taken by residents in the intervention arm will be assessed once using a structured tool to identify opportunities to reduce medication regimen complexity. Residents in the comparison group will receive routine care.</td>
</tr>
<tr>
<td>Key Inclusion and Exclusion Criteria</td>
<td>Key Inclusion Criteria: Permanent residents of aged care facilities who are aged ≥18 years, English-speaking and taking at least one medication. Key Exclusion Criteria: Residents estimated by RACF staff to have less than three months to live and those deemed by facility staff to be medically unstable (e.g. experiencing delirium) will be excluded. Residents may also be excluded at the discretion of RACF staff and their treating clinicians. English-speaking residents who are unable to participate in the Short Assessment of Patient Satisfaction (SAPS) are still eligible for inclusion in the study.</td>
</tr>
<tr>
<td>Study Type</td>
<td>Study type: Interventional Study design: Matched-pair, cluster randomised controlled trial Masking: Open (masking not used) Intervention assignment: Parallel Procedure for enrolling a subject and allocating the treatment (allocation concealment procedures): Study nurses, residents and staff at each RACF will be blinded to the allocation at the time of resident recruitment. Due to the nature of the intervention, it will not be possible to blind study nurses, residents, RACF staff, clinicians and study investigators to the intervention assignment throughout the study period. Methods used to generate the sequence in which subjects will be randomised (sequence generation): Aged care facilities will be paired by the study investigators based on location (i.e. metropolitan or rural) and number of beds. One RACF within each pair will be randomised to the intervention group and the other to the</td>
</tr>
</tbody>
</table>
comparison (usual care) group. An independent pharacoepidemiologist will perform the randomisation using the computerised random number generator within SAS (SAS Institute, Cary, NC). Block randomisation will be used to ensure an equal number of intervention and comparison groups.

<table>
<thead>
<tr>
<th>Date of First Enrollment</th>
<th>24/04/2017</th>
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</thead>
<tbody>
<tr>
<td>Target Sample Size</td>
<td>194 residents</td>
</tr>
<tr>
<td>Recruitment Status</td>
<td>Recruiting</td>
</tr>
</tbody>
</table>

**Primary Outcome(s)**

Primary Outcome: Total number of charted medication administration times over a 24 hour period for regular medications.

Metric/method of measurement: Determined from medication data extracted by the study nurses.

Timepoint: Baseline, and at 4 months after study entry.

**Key Secondary Outcomes**

Secondary Outcome 1: Total number of charted medication administration times over a 24 hour period for regular medications.

Metric/method of measurement: Determined from medication data extracted by the study nurses.

Timepoint: 8 and 12 months after study entry, to assess the sustainability of the intervention.

Secondary outcome 2: Duration of time spent administering medications.

Metric/method of measurement: Determined from medication data extracted by the study nurses and from data collected during a concurrent time-motion study.

Timepoint: Baseline, and at 4 and 8 months after study entry.

Secondary outcome 3: Costs associated with medication administration.

Metric/method of measurement: Determined from medication data extracted by the study nurses and from data collected during a concurrent time-motion study.

Timepoint: Baseline, and at 4 and 8 months after study entry.
| Secondary outcome 4: Resident satisfaction.  
| Metric/method of measurement: Assessed using the 7-item revised version of the Short Assessment of Patient Satisfaction (SAPS) scale.  
| Timepoint: Baseline, and at 4 months after study entry.  

| Secondary outcome 5: Quality of life.  
| Metric/method of measurement: Assessed by a staff informant using the 15-item Quality of Life in Alzheimer’s Disease (QoL-AD) scale adapted for residents of aged care facilities.  
| Timepoint: Baseline, and at 4 months after study entry.  

| Secondary outcome 6: Change in medication incidents (e.g. prescribing errors, pharmacy dispensing errors identified by facility staff, client errors, administration errors or adverse drug reactions).  
| Metric/method of measurement: Determined from the electronic records maintained by the aged care provider organisation.  
| Timepoint: Baseline, and at 4, 8, 12 and 24 months after study entry.  

| Secondary outcome: Number of falls.  
| Metric/method of measurement: Determined from the electronic records maintained by the aged care provider organisation.  
| Timepoint: Baseline, and at 4, 8, 12 and 24 months after study entry.  

| Secondary outcome 8: All-cause overnight hospitalisations.  
| Metric/method of measurement: Determined from the electronic records maintained by the aged care provider organisation.  
| Timepoint: Baseline, and at 4, 8, 12 and 24 months after study entry.  

| Secondary outcome 9: All-cause mortality.  
| Metric/method of measurement: Determined from the electronic records maintained by the aged care provider organisation and/or records maintained by the Government of South Australia Consumer and Business Services: Births, Deaths and Marriages.  

| Timepoint: Baseline, and at 4, 8, 12, 24 and 36 months after study entry. |