Clinical Guideline
Safe Prescribing of New Oral Anticoagulants: Apixaban, Rivaroxaban and Dabigatran

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Summary
New oral anticoagulants (NOAC) may be used as alternatives to warfarin in carefully selected patients. NOAC use is associated with complications such as bleeding, which may range from insignificant to life threatening. The Safe Prescribing of New Oral Anticoagulants: Apixaban, Rivaroxaban and Dabigatran Clinical Guideline assists clinicians in recognising and safely prescribing these agents by taking into account numerous important patient factors with emphasis on renal function. Information about NOAC use in patients undergoing surgical and neuraxial procedures and who may require a temporary interruption to therapy is included.

Keywords
apixaban, rivaroxaban, dabigatran, NOAC, anticoagulant, warfarin, factor Xa thrombin inhibitor, new oral anticoagulant, bleeding, Safe Prescribing of New Oral Anticoagulants

Policy history
Is this a new policy? N
Does this policy amend or update an existing policy? Y
Does this policy replace an existing policy? Y
If so, which policies?
Dabigatran Clinical Guideline: Updated Feb 2014

Applies to
All SA Health Portfolio
All Department for Health and Ageing Divisions
All Health Networks
CALHN, SALHN, NALHN, CHSALHN, WCHN, SAAS Statewide Clinical Services

Staff impact
All Clinical, Medical, Nursing, Allied Health, Emergency, Dental, Mental Health, Pathology, Students

PDS reference
CG220

Version control and change history

<table>
<thead>
<tr>
<th>Version</th>
<th>Date from</th>
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<th>Amendment</th>
</tr>
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<tr>
<td>1.0</td>
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Clinical Guideline: Safe prescribing of new oral anticoagulants: apixaban, rivaroxaban and dabigatran

September 2015
Disclaimer

This guideline provides advice of a general nature. This statewide guideline has been prepared to promote and facilitate standardisation and consistency of practice, using a multidisciplinary approach. The guideline is based on a review of published evidence and expert opinion.

Information in this statewide guideline is current at the time of publication.

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Health practitioners in the South Australian public health sector are expected to review specific details of each patient and professionally assess the applicability of the relevant guideline to that clinical situation.

If for good clinical reasons, a decision is made to depart from the guideline, the responsible clinician must document in the patient’s medical record, the decision made, by whom and detailed reasons for the departure from the guideline.

This statewide guideline does not address all the elements of clinical practice and assumes that the individual clinicians are responsible for:

- discussing care with consumers in an environment that is culturally appropriate and which enables respectful confidential discussion. This includes the use of interpreter services where necessary
- advising consumers of their choice and ensure informed consent is obtained
- providing care within scope of practice, meeting all legislative requirements and maintaining standards of professional conduct, and
- documenting all care in accordance with mandatory and local requirements.

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Clinical Guideline: Safe prescribing of new oral anticoagulants: apixaban, rivaroxaban and dabigatran v1.1

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Introduction

Apixaban, rivaroxaban and dabigatran are ‘new’ or ‘novel’ oral anticoagulant medicines (often referred to as NOAC). They are listed on the Australian Pharmaceutical Benefits Scheme (PBS) for treatment and prevention of thrombo-embolic disease. NOAC are also available on the South Australian Medicines Formulary (SAMF) for specific indications.

Background

Safe and effective use of any anticoagulant requires careful patient selection and clinical monitoring to minimise the risk of thrombosis and of bleeding.

Prescribing a NOAC requires an in-depth knowledge of their pharmacology and clinical use, careful patient selection and monitoring to ensure the best outcomes.

The NOAC are different to existing oral anticoagulants, in particular warfarin, in their clinical monitoring requirements, drug interactions and the limited options available for reversal.

Bleeding complications with NOAC are potentially severe and have been fatal in an often fragile population.

Serious incidents involving NOAC use have been reported due to:

- clinicians not recognising these medications as anticoagulants
- concurrent prescribing of other anticoagulant medications e.g. heparin
- misinterpretation of coagulation tests and monitoring requirements
- inappropriate or premature cessation prior to procedures
- drug interactions.

This guideline has been developed to assist SA Health staff to safely use NOAC by:

- recognising the medicines by generic and trade name
- managing the risk of thrombosis versus the bleeding risk
- considering individual patient bleeding risk factors
- understanding the significance of different coagulation test results and drug interactions
- making appropriate decisions regarding surgery and neuraxial procedures.

A Clinical Guideline: Management of bleeding associated with apixaban, rivaroxaban and dabigatran is also available.

These guidelines are based on evidence available at the time and do not replace expert medical judgement. For more comprehensive guidance please refer to a local haematologist and the manufacturer’s product information.
Standards

The information in this guideline aligns with the Australian Commission on Safety and Quality in Health Care - National Safety and Quality Health Service Standards: Standard 4 - Medication Safety.¹

> Action item 4.11 - Identifying high-risk medicines in the organisation and ensuring they are stored, prescribed, dispensed and administered safely.
General

1. South Australian Medicines Formulary Recommendations

1.1. Apixaban (Eliquis®) - preferred NOAC
   - 2.5 mg and 5 mg tablets for the prevention of stroke or systemic embolism in patients with non-valvular atrial fibrillation (AF) as per PBS criteria.\(^3\)
   - 2.5 mg tablets for the prevention of venous thromboembolism in total hip or knee replacement as per PBS criteria.\(^3\)

1.2. Rivaroxaban (Xarelto®) - restricted inclusion
   - 15 mg and 20 mg tablets for initial and continuing treatment of deep vein thrombosis (DVT) and pulmonary embolism (PE) as per PBS criteria.\(^3\)
   - 10 mg tablets for the prevention of venous thromboembolism in total hip or knee replacement as per PBS criteria.\(^3\)

   Rivaroxaban is not listed on the South Australian Medicines Formulary (SAMF) for initiation of therapy for prevention of systemic embolism or stroke in non-valvular AF.

1.3. Dabigatran (Pradaxa®) - not listed on SAMF, however:
   - 150 mg capsules are available via a streamlined Individual Patient Use (IPU) process for initiation of the prevention of stroke or systemic embolism in patients who:
     - fulfil the PBS criteria\(^3\) with non-valvular atrial fibrillation and have one or more risk factors for developing stroke or systemic embolism AND
     - < 75 years
     - creatinine clearance (Cockcroft and Gault calculation) > 50 mL / min
     - CHADS\(_2\) score ≥ 2
     - not taking a strong P- glycoprotein inhibitor
     - does not have active gastrointestinal (GI) disease or a high risk of GI bleeding
     - able to swallow capsules whole
     - does not require a dosage administration aid.

2. Pharmacological Characteristics of NOAC

Table 1: Pharmacological Characteristics of NOAC \(^4,5\)

<table>
<thead>
<tr>
<th>Apixaban (Eliquis®)</th>
<th>Rivaroxaban (Xarelto®)</th>
<th>Dabigatran etexilate (prodrug of dabigatran) (Pradaxa®)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Mechanism of action</strong></td>
<td>Direct Factor Xa inhibitor</td>
<td>Direct Factor Xa inhibitor</td>
</tr>
<tr>
<td><strong>T(_{max})</strong></td>
<td>1 – 3 hours</td>
<td>2.5 – 4 hours</td>
</tr>
<tr>
<td><strong>Half-life (hours) if CrCl is 30 - 50 mL/min.</strong></td>
<td>8 - 15</td>
<td>5 - 9 (healthy) 11 - 13 (elderly)</td>
</tr>
<tr>
<td><strong>Elimination</strong></td>
<td>Renal 27% (multiple other routes)</td>
<td>Renal 33% Renal metabolites 33% Hepatic 33%</td>
</tr>
</tbody>
</table>
3. Principles for prescribing NOAC

> Prescribing a NOAC requires an in depth knowledge of their pharmacology and clinical use. Careful patient selection and monitoring ensure the best outcomes.
> A risk-benefit assessment should always be conducted prior to prescribing NOAC for any patient.
> It is essential that creatinine clearance is calculated (using the Cockcroft-Gault equation and ideal body weight) and documented before prescribing a NOAC.

Appendix 1: ‘Flowchart for Prescribing New Oral Anticoagulants Apixaban, Rivaroxaban and Dabigatran’ provides a summary of the risk factors that must be taken into account when considering initiation treatment with these medications.

> Drug accumulation and excessive anticoagulation can occur, especially if renal function is impaired. Renal function must be monitored at least annually. Monitoring should be increased to 3 or 6 monthly when other factors may indicate declining renal function or dehydration.

4. Drug Interactions

NOAC have fewer drug interactions than warfarin, however many clinically significant interactions exist. Individual patient bleeding risks must be considered and specialist advice sought as these are often complex situations.

The European Heart Rhythm Association’s ‘Practical Guide on the use of new oral anticoagulants in patients with non-valvular atrial fibrillation’ provides a useful drug interaction chart that includes advice on NOAC dosage adjustments. The Australian Medicines Handbook (AMH), consumer product information sheets and NPS Medicinewise (for health professionals, decision making tools) website also provide comprehensive lists of interactions. Note that some recommendations may differ.

Renal function should be rechecked if drugs that may impair renal function are commenced. For example, non-steroidal anti-inflammatory drugs, diuretics or angiotensin-converting enzyme (ACE) inhibitors. These combinations should be avoided wherever possible.

4.1. Thrombolytic agents or antiplatelet agents (e.g. aspirin, clopidogrel, prasugrel, ticagrelor, ticlopidine):

Concomitant administration of these agents with NOAC is not recommended unless clinically indicated because dual therapy is associated with higher risks of bleeding and anaemia, especially if there are other risk factors. Seek specialist advice.

4.2. Other anticoagulants:

Concomitant administration with NOAC is not recommended. This includes bridging therapy as NOAC have a relatively rapid onset of action.

4.3. Non-steroidal anti-inflammatory drugs (NSAIDs) and Cox 2 inhibitors:

Monitor for risks of bleeding if a NOAC is used with NSAIDs, especially those with half-lives greater than 12 hours (for example, naproxen, piroxicam). This combination should be avoided due to the increased risk of gastrointestinal bleeding.

4.4. Strong inhibitors of CYP3A4 or P-glycoprotein (P-gp) transporter:

All three NOAC are substrates for the P-glycoprotein transporter. Apixaban and rivaroxaban are also metabolised largely by CYP3A4. Inhibitors of these systems increase NOAC bioavailability. See flowchart, Appendix 1 and Table 2a for a list of common drugs affected.

NOAC are contraindicated with drugs that are strong inhibitors of both of these pathways including:

> systemic azole antifungals: ketoconazole, itraconazole, voriconazole, and posaconazole.
> HIV protease inhibitors: for example, ritonavir, saquinavir.
> dronedarone
4.5. **Less potent inhibitors or drugs inhibiting only one of these two pathways**
These drugs may significantly contribute to increased bleeding risk if administered with apixaban or rivaroxaban when there are other bleeding risk factors; however, they are not currently contraindicated. A dose reduction may be appropriate and concomitant administration should be considered on an individual basis.

- cardiac medicines – consider cardiology advice
  - calcium channel blockers; for example, diltiazem, verapamil
  - quinidine
  - amiodarone
- fluconazole
- immunosuppressants; for example, cyclosporine, tacrolimus
- macrolides; for example, clarithromycin, erythromycin
- others; for example, dipyridamole, tamoxifen.

4.6. **Strong inducers of CYP3A4:**
Concomitant use of a NOAC with strong P-gp or CYP3A4 inducers may lead to reduced plasma levels of the NOAC and an increase in stroke risk. Examples of strong inducers of CYP3A4 include:

- rifampicin, phenytoin, carbamazepine, phenobarbitone or St John’s Wort

The combination with apixaban or dabigatran is contraindicated.
If using rivaroxaban in these situations, a dose change should be considered but the combination is preferably avoided as there is no way to monitor the effectiveness of the NOAC.

4.7. **Contraindicated drug combinations**
Avoid concomitant use due to clinically significant increased plasma levels and increased bleeding risk.

*Table 2a: Contraindicated drug combinations with NOAC*[^4][^5]

<table>
<thead>
<tr>
<th>Apixaban, Rivaroxaban and Dabigatran</th>
<th>Dabigatran</th>
</tr>
</thead>
<tbody>
<tr>
<td>Systemic azole antifungals:</td>
<td>Simultaneous initiation with verapamil</td>
</tr>
<tr>
<td>&gt; ketoconazole, itraconazole, voriconazole, posaconazole (except fluconazole)</td>
<td></td>
</tr>
<tr>
<td>HIV-protease inhibitors:</td>
<td></td>
</tr>
<tr>
<td>&gt; for example, ritonavir, saquinavir</td>
<td></td>
</tr>
<tr>
<td>Dronedarone</td>
<td></td>
</tr>
</tbody>
</table>

4.8. **Drug combinations that are preferably avoided**
Consider alternative therapy due to increased bleeding risk, or seek advice as dose change may be warranted.
The presence of renal or hepatic impairment may make these interactions significant, even with weak inhibitors.
During concomitant use, monitor patients closely for bleeding and encourage them to report signs of bleeding.
Table 2b. Drug combinations that are preferably avoided with all NOACs:

<table>
<thead>
<tr>
<th>Apixaban and Rivaroxaban and Dabigatran</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Increased plasma level of NOAC and increased bleeding risk</strong></td>
</tr>
<tr>
<td><strong>Less potent P-gp and/or CYP3A4 inhibitors:</strong></td>
</tr>
<tr>
<td>&gt; cardiac medications – <em>consider cardiology advice</em></td>
</tr>
<tr>
<td>&gt; amiodarone, diltiazem, quinidine, verapamil – especially simultaneous initiation</td>
</tr>
<tr>
<td>&gt; clarithromycin, erythromycin</td>
</tr>
<tr>
<td>&gt; cyclosporin, tacrolimus</td>
</tr>
<tr>
<td>&gt; fluconazole</td>
</tr>
<tr>
<td>&gt; grapefruit juice.</td>
</tr>
</tbody>
</table>

| **Reduced plasma levels of NOAC and increased thromboembolic risk** |
| **Potent P-gp/CYP3A4 inducers:** |
| > carbamazepine, phenobarbitone, phenytoin, rifampicin, St John’s Wort |

| **Pharmacodynamic interactions that increase bleeding risk** |
| > antiplatelet / anticoagulant / antithrombotic agents: |
| > for example, aspirin, cox-2 inhibitors, fish oil, clopidogrel, prasugrel, ticagrelor, heparins, warfarin, alteplase |
| > systemic steroid therapy |
| > chemotherapy |

| **Some antidepressants:** |
| > for example, selective serotonin (SSRI) or serotonin-noradrenaline (SNRI) reuptake inhibitors |

5. Transitions of care

The following factors must be communicated through all transitions of care:

> Monitor renal function at least annually, or when other patient factors such as dehydration or declining renal function suggest that an increased frequency of monitoring is necessary. For example, in older people (≥ 75 years) or when CrCl < 50 mL/min, monitoring every 3–6 months may be warranted.

> Declining renal function or concomitant use of agents that affect renal function are important factors in increasing the bleeding risk associated with NOAC.

> Any planned dose change, or expected limits on duration of treatment must be clearly documented. For example with rivaroxaban when used for the treatment and subsequent prevention of acute and recurrent DVT or PE.

> Patient counselling should include:

  > communicating the discharge from hospital care plan with patient / carer
  > encouraging regular and adequate follow up
  > educating the patient and carer, including information on how to recognise and respond to bleeding by seeking medical attention immediately if bleeding is suspected.
6. Laboratory Tests

- Regular routine monitoring of International Normalised Ration (INR) or other coagulation parameters is **not** necessary.
- The anticoagulant effect of the NOAC should be measured if:
  - the patient is bleeding severely
  - urgent surgery is required
  - a thrombotic event occurs or there is a recurrence or exacerbation of thromboembolism.
- The time of the last NOAC dose is required to interpret the results.
- Interpretation of coagulation tests must take into account the clinical setting.
- When the results of coagulation test results are prolonged, extend testing to confirm the likelihood of a drug effect with Hemoclot or anti-Xa, and refer to Table 3.
- All ‘standard’ coagulation assays (PT, aPTT, TT) may be normal in the presence of significant levels of apixaban.
- Specific assays to assess drug level (anti-Xa assay for rivaroxaban and apixaban, Hemoclot for Dabigatran) are not available at all sites or 24 hours a day. If needed urgently, consult with the on-call clinical Haematology Service.

**Table 3: Effect of New Oral Anticoagulants on Laboratory Coagulation Tests**

<table>
<thead>
<tr>
<th>Tests</th>
<th>Apixaban</th>
<th>Rivaroxaban</th>
<th>Dabigatran</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prothrombin time (PT) and</td>
<td>Insensitive</td>
<td>Relatively insensitive</td>
<td>Insensitive</td>
</tr>
<tr>
<td>International Normalised Ratio (INR)</td>
<td>PT and INR not recommended</td>
<td>A 1.5 to 2.0 fold increase will be seen at peak concentrations</td>
<td>A prolonged PT in the absence of other causes often indicates excess drug levels</td>
</tr>
<tr>
<td>Activated Partial Thromboplastin Time (aPTT)</td>
<td>Insensitive</td>
<td>aPTT is prolonged dose dependently, but is less sensitive than PT</td>
<td>Variable sensitivity A normal aPTT excludes high drug levels</td>
</tr>
<tr>
<td>Thrombin Time (TT)</td>
<td>Insensitive</td>
<td>Insensitive</td>
<td>Very sensitive. A normal TT excludes the presence of Dabigatran</td>
</tr>
<tr>
<td>Chromogenic anti-Xa assay</td>
<td>Use modified anti-Xa assay for apixaban</td>
<td>Use modified anti-Xa assay for rivaroxaban</td>
<td>Insensitive</td>
</tr>
<tr>
<td>Hemoclot (Not always available)</td>
<td>Not suitable</td>
<td>Not suitable</td>
<td>Can quantify drug level</td>
</tr>
</tbody>
</table>
Protocols

1. Total Hip and Knee Replacement

Haemostasis must be established prior to initiating NOAC.

Table 4: Dose for prevention of venous thromboembolism (VTE) for total hip or total knee replacement

<table>
<thead>
<tr>
<th>Creatinine clearance</th>
<th>Apixaban</th>
<th>Rivaroxaban</th>
</tr>
</thead>
<tbody>
<tr>
<td>CrCl &gt; 50 mL/min</td>
<td>SAMF preferred NOAC for this indication</td>
<td>10 mg once daily</td>
</tr>
<tr>
<td>CrCl 30 - 50 mL/min</td>
<td>2.5 mg twice daily</td>
<td></td>
</tr>
<tr>
<td>CrCl 25 - 29 mL/min</td>
<td></td>
<td>contraindicated</td>
</tr>
<tr>
<td>CrCl &lt; 25 mL/min</td>
<td>contraindicated</td>
<td></td>
</tr>
</tbody>
</table>

> Duration of therapy:
  > Hip - up to 35 days
  > Knee - up to 15 days

Table 5: Time of the first dose of NOAC following total hip or knee replacement

<table>
<thead>
<tr>
<th>Time elapsing between surgery and the first dose after surgery</th>
<th>Apixaban</th>
<th>Rivaroxaban</th>
</tr>
</thead>
<tbody>
<tr>
<td>12 – 24 hours</td>
<td>6 – 10 hours</td>
<td></td>
</tr>
</tbody>
</table>

For other surgery and use of NOAC for other indications refer to Tables 8 and 9.

2. Deep vein thrombosis, pulmonary embolism and atrial fibrillation

> In general the risk-benefit balance in AF and PE favour the use of an anticoagulant regardless of age and renal function. Both the stroke risk and the bleeding risk increase with age, and renal function declines. Being over 75 years of age, declining renal function and other patient factors may increase the bleeding risk and outweigh benefit of NOAC therapy.

> It is recommended that prescribers measure renal function and consider this with the patient’s bleeding risk (HASBLED score) and stroke risk (CHADSVASC score). NOAC administration should not necessarily be avoided in favour of warfarin (or no treatment) if renal function is reduced, and where CrCl ≥ 30 mL / min. Seek specialist advice in complex situations.
2.a. Acute and Recurrent Deep Vein Thrombosis or Pulmonary Embolism

Rivaroxaban is currently the only NOAC approved by the TGA and PBS listed for the initial and continuing treatment of venous thromboembolism and pulmonary embolism. Rivaroxaban prescribing within SA Health is currently restricted to this indication.

NOTE: In March 2015, the Pharmaceutical Benefits Advisory Committee (PBAC) recommended the listing of apixaban 2.5mg bd or 5mg bd for the treatment of acute and recurrent DVT or PE; however, at the time of printing this guideline apixaban has not been added to the PBS for these indications. (PBAC Outcomes March 2015)

Table 6: Treatment or prevention of acute and recurrent DVT or PE

<table>
<thead>
<tr>
<th>Creatinine clearance</th>
<th>Rivaroxaban 15 and 20 mg*</th>
</tr>
</thead>
<tbody>
<tr>
<td>CrCl ≥ 30 mL/min</td>
<td>15 mg twice daily with food for 3 weeks, followed by 20 mg once daily, for a minimum of 3 months.</td>
</tr>
<tr>
<td></td>
<td>The decision to extend treatment past 3 months should be based on the risk of recurrent thrombosis and haematology referral is suggested if the risk/benefit of continued anticoagulation is unclear.</td>
</tr>
<tr>
<td></td>
<td>Current experience for more than 12 months is limited</td>
</tr>
<tr>
<td>CrCl &lt; 30 mL/min</td>
<td>Contraindicated</td>
</tr>
</tbody>
</table>

* Rivaroxaban must be taken with food as there is increased absorption from 66% in fasted state to nearly 100% with food.

2.b. Non-valvular Atrial Fibrillation

Rivaroxaban is not listed on the SA Health Medicines Formulary for initiation of therapy for prevention of systemic embolism or stroke in non-valvular AF.

Table 7: Dose for non-valvular atrial fibrillation

<table>
<thead>
<tr>
<th>Creatinine clearance</th>
<th>Apixaban (SAMF preferred)</th>
<th>Rivaroxaban* (SAMF: not available for initiation of therapy)</th>
<th>Dabigatran (SAMF: 150 mg only is available as IPU)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CrCl &gt; 50 mL/min</td>
<td>5 mg twice daily or if any 2 of the following are present (even if CrCl &gt; 25 mL/min: older than 80 years weight ≤ 60 kg serum creatinine ≥133 micromol / L then use: 2.5 mg twice a day</td>
<td>20 mg once daily</td>
<td>150 mg twice daily not recommended by SAMF if there are bleeding risk factors other than renal function</td>
</tr>
<tr>
<td>CrCl 31 - 50 mL/min</td>
<td>&gt; older than 80 years &gt; weight ≤ 60 kg</td>
<td>15 mg once daily</td>
<td>not recommended</td>
</tr>
<tr>
<td>CrCl 25 - 30 mL/min</td>
<td>&gt; serum creatinine ≥133 micromol / L then use: 2.5 mg twice a day</td>
<td>contraindicated</td>
<td>contraindicated</td>
</tr>
</tbody>
</table>

* Rivaroxaban must be taken with food as there is increased absorption from 66% in fasted state to nearly 100% with food.
3. Surgery

> Consider the risk of thrombotic complications in the perioperative period if the NOAC is stopped, relative to the risk of bleeding if it is continued.
> Discontinuing the NOAC may not be essential, especially with very minor procedures.
> Time the surgery to coincide with minimal residual anticoagulant effect as per the elimination half-life of the NOAC in relation to the patient’s renal function, based on calculated creatinine clearance, and the bleeding risk.

3.a. Urgent Surgery

> If possible delay surgery until coagulation screen is normal or sufficient time has passed for drug clearance.
> Seek haematologist’s advice if surgery cannot be sufficiently delayed.

3.b. Pre-operative interruption of NOAC

Table 8: Suggested time between last dose of NOAC and surgery

<table>
<thead>
<tr>
<th>Renal Function and related half-life*</th>
<th>NOAC and current dose</th>
<th>Low bleeding risk† surgery (2 to 3 half-lives between last dose and surgery)</th>
<th>Time of last dose before surgery (hours)</th>
<th>High bleeding risk‡ surgery (4 to 5 half-lives between last dose and surgery)</th>
<th>Time of last dose before surgery (hours)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CrCl ≥ 50 mL/min (half-life 7 - 8 hours)</td>
<td>apixaban 5mg twice daily</td>
<td></td>
<td>24</td>
<td>48 - 72</td>
<td></td>
</tr>
<tr>
<td>CrCl 30 - 49 mL/min (half-life 17 - 18 hours)</td>
<td>apixaban 5mg twice daily</td>
<td></td>
<td>48</td>
<td>72</td>
<td></td>
</tr>
<tr>
<td>CrCl ≥ 50 mL/min (half-life 5 - 9 hours)</td>
<td>rivaroxaban 20mg once daily</td>
<td></td>
<td>24</td>
<td>48 - 72</td>
<td></td>
</tr>
<tr>
<td>CrCl 30 - 49 mL/min (half-life 9 - 13 hours)</td>
<td>rivaroxaban 20mg once daily</td>
<td></td>
<td>48</td>
<td>72</td>
<td></td>
</tr>
<tr>
<td>CrCl ≥ 50 mL/min (half-life 12 - 17 hours)</td>
<td>dabigatran 150 mg twice daily</td>
<td></td>
<td>24</td>
<td>48 - 72</td>
<td></td>
</tr>
<tr>
<td>CrCl 30 - 49 mL/min (half-life 13 - 23 hours)</td>
<td>dabigatran 150 mg twice daily</td>
<td></td>
<td>48 - 72</td>
<td>96</td>
<td></td>
</tr>
</tbody>
</table>

* Estimate of half-life is based on calculated renal clearance using the Cockcroft-Gault equation
† Aim for mild to moderate residual anticoagulant effect at surgery (<12-25%)
‡ Aim for no or minimal residual anticoagulant effect at surgery (<3-6%)
3.c. Post-operative NOAC recommencement

Ensure that renal and hepatic functions are normal before restarting NOAC.

Time the resumption of a NOAC based on the anticipated surgical bleeding risk, the extent of intra-operative or post-operative bleeding and the patient’s risk of thromboembolism.

In patients undergoing a procedure associated with high bleeding risk, the resumption of therapeutic anticoagulation with a NOAC should normally be delayed 48 to 72 hrs. Appropriate anticoagulant therapy should be administered during this time as indicated for the procedure. Management of patients at high risk of thrombotic complications (see local guidelines) should be guided by a consultant, and refer to Table 12 for converting the patient to NOAC.

Consider intermittent pneumatic compression if prophylactic anticoagulation cannot be safely administered.

Table 9: Suggested time between surgery and resumption of NOAC\(^4,5\)

(For timing of first NOAC dose following total hip or knee replacement, refer to Table 5).

<table>
<thead>
<tr>
<th>NOAC*</th>
<th>Low bleeding risk</th>
<th>High bleeding risk</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Resume 24 hours after surgery</td>
<td>Resume 48 -72 hours after surgery</td>
</tr>
</tbody>
</table>

* For patients at high risk of thromboembolism consider commencing NOAC sooner, but with a reduced dose, in the evening after surgery and on the following day:

- dabigatran: 75 mg once daily
- rivaroxaban: 10 mg once daily
- apixaban: 2.5 mg twice daily for 2 days or until it is safe to resume therapeutic anticoagulation.

For patients at high risk of thromboembolism consider commencing NOAC sooner, but with a reduced dose, in the evening after surgery and on the following day:

- dabigatran: 75 mg once daily
- rivaroxaban: 10 mg once daily
- apixaban: 2.5 mg twice daily for 2 days or until it is safe to resume therapeutic anticoagulation.
4. Neuraxial anaesthesia (or lumbar puncture)

- A patient is at risk of developing an epidural or spinal haematoma if a neuraxial procedure is undertaken when anticoagulated.
- There is a lack of published evidence regarding the safety of neuraxial anaesthesia in patients therapeutically anticoagulated with NOAC.
- Avoid neuraxial procedures until laboratory testing (if available) establishes the absence of any anticoagulant effect or wait until five renally adjusted half-lives have elapsed since the last NOAC dose.
- Always monitor carefully for signs and symptoms of neurological impairment.
- Seek specialist advice from Haematology, Anaesthesia or Acute Pain Service.

5. Switching to or from other anticoagulants

5.a. Switching from warfarin to a NOAC

When deciding whether to transition a patient from warfarin to a NOAC the following should be considered:

- patients stable on warfarin (ie INR is within range for > 65% of time in 3 months) should continue on warfarin
- patient preference
- lack of reliable agents to reverse severe NOAC related bleeding
- access to INR testing versus the added convenience of NOAC therapy
- the potential for enhanced efficacy and the reduced risk of intracranial bleeding with NOAC, as compared with warfarin in correctly selected patients
- other risk factors such as drug interactions and consistency of dietary vitamin K intake.

Recommended strategy:
- Discontinue warfarin and start NOAC when INR ≤ 2.5.

5.b. Switching from a NOAC to warfarin

When converting a patient from NOAC to warfarin it is necessary to consider the following:

- the elimination half-life of the NOAC is affected by renal function
- there is typically a 5-day delay in the onset of warfarin effect
- the INR may be affected by both NOAC and warfarin, hence the INR will better reflect warfarin only after the NOAC has been stopped for at least two days.

Recommended strategy:
- Start daily warfarin dose of ≤ 5mg. Higher initiation doses are not recommended.5
- Continue NOAC
- Measure the first INR on day 2 or 3 after initiating warfarin. The main purpose is to identify high levels of warfarin and maintain caution with ongoing doses.
- Do not use point of care monitors during the changeover period between NOAC and warfarin.
- Stop NOAC when INR ≥ 2 on two consecutive days, taking into account the NOAC effect on INR.
- Consult haematology services for further advice.
### Table 10: Recommended time between initiating warfarin and ceasing NOAC\(^5\)

<table>
<thead>
<tr>
<th>Calculated creatinine clearance</th>
<th>Number of days after initiating warfarin that <strong>apixaban or rivaroxaban</strong> should be stopped</th>
<th>Number of days after initiating warfarin that <strong>dabigatran</strong> should be stopped</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt; 50 mL/min</td>
<td>4 days</td>
<td>3 days</td>
</tr>
<tr>
<td>31 to 50 mL/min</td>
<td>3 days</td>
<td>2 days</td>
</tr>
<tr>
<td>15 to 30 mL/min</td>
<td>2 days</td>
<td>1 day</td>
</tr>
<tr>
<td>&lt; 15 mL/min</td>
<td>Consult haematology service</td>
<td></td>
</tr>
</tbody>
</table>

### Table 11: Converting from injected anticoagulants to NOAC\(^5\)

<table>
<thead>
<tr>
<th>Converting from</th>
<th>Converting to</th>
<th>Instructions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low molecular weight heparin (LMWH)</td>
<td>NOAC</td>
<td>Discontinue LMWH and start NOAC when the next dose of LMWH would have been due</td>
</tr>
<tr>
<td>Continuous heparin infusion</td>
<td>NOAC</td>
<td>Discontinue the infusion and start NOAC immediately</td>
</tr>
</tbody>
</table>

### Table 12: Converting from NOAC to injected anticoagulants\(^5\)

<table>
<thead>
<tr>
<th>Converting from</th>
<th>Converting to</th>
<th>Instructions</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>apixaban or rivaroxaban</strong></td>
<td>Low molecular weight heparin (LMWH) or unfractionated heparin</td>
<td>Discontinue NOAC and start LMWH or heparin 12-24 hours after the last dose of NOAC. A bolus dose of unfractionated heparin is not required</td>
</tr>
<tr>
<td><strong>dabigatran</strong></td>
<td>LMWH</td>
<td>If CrCl ≥ 30 mL/min: &gt; discontinue dabigatran and start LMWH 12 - 24 hours after last dose of dabigatran. If CrCl &lt; 30 mL/min: &gt; LMWH is not recommended and dabigatran is contraindicated</td>
</tr>
<tr>
<td><strong>dabigatran</strong></td>
<td>unfractionated heparin</td>
<td>If CrCl ≥ 30 mL/min: &gt; discontinue dabigatran and start unfractionated heparin 12 - 24 hours after last dose of dabigatran If CrCl &lt; 30 mL/min: &gt; discontinue dabigatran and start unfractionated heparin 48 hours after the last dose of dabigatran.</td>
</tr>
</tbody>
</table>
Acknowledgements / Consultation

SA Health thanks all staff involved in the preparation and review of this material, in particular:

- **NOAC Working group:**
  - Dr Simon McRae (Director Haemophilia Treatment Centre, RAH)
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  - Mr Richard Marotti (Pharmacy Director, LMH)
  - Mr Vaughn Eaton (Pharmacy Director, FMC)
  - Ms Naomi Burgess (Director, MTPP)
  - Ms Elizabeth Campbell (Project Pharmacist, MTPP)

- Dr W. Heddle, Arrhythmia Working Group, Cardiology Clinical Network

- Ms Kaye Barratt, Senior Specialist Pharmacist, Medicines and Technology Policy and Programs

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Appendices

1. **Flowchart for Prescribing New Oral Anticoagulants Apixaban, Rivaroxaban and Dabigatran.**

   This flowchart summarises the factors that must be taken into account when considering the initiation of treatment with apixaban, rivaroxaban or dabigatran.
Flowchart for Prescribing New Oral Anticoagulants (NOAC) Apixaban, Rivaroxaban and Dabigatran

Calculate and record creatinine clearance (CrCl)
(use Cockcroft - Gault equation)
Record full blood count and liver function

Take a detailed history
Check all laboratory considerations and exclusion criteria

Assess bleeding risk

Consider concomitant medicines

If the patient is on warfarin
and
if all other patient factors warrant the changeover to a NOAC then stop warfarin and see guideline instructions for converting patient from warfarin to NOAC

Laboratory considerations

Renal function
- rivaroxaban is contraindicated if: CrCl < 30 mL/min
- apixaban is contraindicated if: CrCl < 25 mL/min
- dabigatran is contraindicated in SA Health, for initiation of therapy if: CrCl < 50 mL/min (see dabigatran below)

Liver disease
Contraindicated if alanine transaminase (ALT) > 2 x upper limit of normal,
or for apixaban Child-Pugh C (if B use with caution)
or rivaroxaban and dabigatran Child-Pugh B and C.

Full Blood Count
Anaemia Hb ≤ 100 g/L

Assess bleeding risk (seek specialist advice if ‘yes’ to any of the following):
- history of significant bleeding
- surgery ≤ 1 month ago
- gastro-intestinal (GI) bleed ≤ 12 months ago
- Gl ulcer ≤ 30 days ago
- fibrinolytic treatment ≤ 24 hours ago
- on any anticoagulation agent
- on dual antiplatelet therapy
- platelet count < 100 x 10^9 /L

Apixaban (Eliquis®)
Total hip or knee replacement (VTE prophylaxis) 2.5 mg twice a day
hip: up to 35 days / knee: up to 15 days
Non-valvular AF 5 mg twice a day
or
If any 2 of the following are present:
- age ≥ 80 years,
- weight ≤ 60 kg or
- serum creatinine ≥ 133 micromol/L
2.5 mg twice daily

Rivaroxaban (Xarelto®)
SAMF restricted to:
Total hip or knee replacement (VTE prophylaxis) 10 mg once daily
hip: up to 35 days / knee: up to 15 days
Initial and continuing treatment of deep vein thrombosis (DVT) and pulmonary embolism (PE)
(if CrCl > 30 mL/min)
15 mg twice daily for 3 weeks,
then reduce to 20 mg daily

Dabigatran (Pradaxa®)
Streamlined Individual Patient Use Authority for:
Non-valvular AF 150 mg twice daily only in selected patients
(if CrCl ≥ 50 mL/min)
also refer to SA Medicines Formulary

Exclusion criteria
- < 18 years
- known hypersensitivity to NOAC
- pregnant or breastfeeding
- active significant bleeding or disorder of haemostasis (von Willebrand’s or coagulation deficiency)
- prostatic heart valve or severe valvular disease
- recent stroke – relative contraindication (seek specialist advice)
- thrombus and recent stent (seek cardiologist advice)
- active cancer – relative risk (seek specialist advice)

This is not an exhaustive list – refer to guideline. The European Heart Rhythm Association provides a useful decision making chart.

Concomitant medicines
Contraindicated:
- Potent P-glycoprotein (P-gp) competitors and CYP3A4 inhibitors:
  - ketoconazole, itraconazole, posaconazole, voriconazole
  - HIV protease inhibitors e.g. ritonavir, saquinavir
  - dronedarone
- Enzyme inducers: contraindicated with apixaban and dabigatran e.g. rifampicin, St John’s Wort, carbamazepine, phenytoin, and phenobarbitone. Preferably avoid with rivaroxaban.

Preferably avoided: known or expected increases in NOAC blood levels may occur with the following medicines and a NOAC dose reduction may be appropriate; consider on an individual basis:
- Cardiac medicines – consider cardiologist advice
  - verapamil, especially simultaneous initiation (formulations differ)
  - quinidine
  - amiodarone
- fluconazole
- cyclosporin, tacrolimus
- erythromycin, clarithromycin
- If antiplatelet, anticoagulant or antithrombotic agents are required seek haematologist advice
References

1. Australian Commission on Safety and Quality in Health Care - National Safety and Quality Health Service Standards: Standard 4 - Medication Safety
2. South Australian Medicines Formulary (SAMF)

Related documents and references

1. Prescribing information – MIMS ONLINE
2. Apixaban (Eliquis), dabigatran (Pradaxa) and rivaroxaban (Xarelto): Information for health professionals (TGA, All alerts)
3. Dabigatran (Pradaxa) and the risk of bleeding: Information for health professionals (TGA, All alerts)
4. NPS Medicinewise (For health professionals)
5. Newer Oral Anticoagulants (Update); New South Wales Health, Safety notice 002/14
6. European Society of Cardiology Guidelines

Patient information:

1. Living with a new Oral Anticoagulant NOAC (WATAG) - patient guideline 2013
2. NPS Medicinewise
3. Dabigatran (Pradaxa) safety update: Information for consumers; (TGA, All alerts)
4. Apixaban (Eliquis), dabigatran (Pradaxa) and rivaroxaban (Xarelto): Information for consumers (TGA, All alerts)