Anticoagulation with Warfarin

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**Introduction:**
Many adults with congenital heart defects need to take oral anticoagulants. Common reasons to need this type of treatment include prosthetic heart valve replacements, atrial arrhythmias, and previous surgeries such as the Fontan operation. Other diagnoses include conditions where there is blood flow that is turbulent caused by narrow vessels, aneurysms, poor function or artificial devices. Anticoagulants slow the time it takes to form a blood clot. These medications also prevent complications like valve obstruction, blood clots forming on artificial valves, and blood clots traveling to the pulmonary bed causing a pulmonary embolism or to the brain causing a stroke.

**Lesions:** (Monagle, 2004, 2008)

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<th>Indications/Goal Ranges</th>
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<tr>
<td>Indication</td>
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<td>Prosthetic heart valves – Aortic position</td>
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**Critical Thinking Points:**
- Initial Response is usually with 24 hours after administration
- Peak anticoagulation response 72 to 96 hr
- Duration of action of a single dose is 2-5 days
- Half life: Following a single warfarin dose, the terminal half-life is about 1 week with a mean effective half-life of 40 hr (range, 20 to 60 hr).

*(Prod Info COUMADIN(R) oral tablets, intravenous injection, 2010)*
**Diagnostic Evaluation:**
- International Normalized Ratio (INR) means that no matter how or where the lab is drawn it can be compared to another INR. The INR is a measure of how long it takes the body to form a clot.
  - All people, regardless of anticoagulation status, have a baseline INR around 0.8-1.0. An elevated baseline INR should be investigated before beginning anticoagulation therapy
- Many centers draw a baseline INR and CBC before an initial dose of an anticoagulant medication is given.

**Treatment:**
- Initial dose:
  - Loading dose is generally unnecessary.
    - Theoretical reasons for beginning treatment with the average maintenance dose of ≈5 mg daily
    - Usually results in an INR of ≥ 2.0 after 4 or 5 days.
  - When a rapid effect is required, heparin or low molecular weight heparin should be given concurrently with warfarin for 4 days.
  - Loading dose:
    - Warfarin dosing may be separated into initial and maintenance phases
    - INR response is monitored frequently until a stable dose-response relationship is obtained.
    - Once the INR is stable, the frequency of INR testing is reduced.
    - Anticoagulant effect is observed within 2 to 7 days after beginning oral warfarin.
- Guidelines when using a Loading dose (Children’s Hospital Boston Formulary, Boston, Massachusetts):
  - Loading Dose Day 1
    1. Give 0.1 mg/kg (maximum 0.2 mg/kg) up to a maximum of 15 mg per dose
      a. The 0.1mg/kg dose is a routine dose.
      b. The higher end of loading dose (0.2mg/kg) may be considered when prompt elevation in INR desired and the patient’s INR can be monitored very closely (usually on a daily basis). Most often this loading dose is used in patients who have had cardiac surgery for implantation of mechanical heart valves.
  2. A loading dose lower than 0.1mg/kg should be considered in patients with Fontan physiology, liver disease, decreased vitamin K intake and/or who are known slow metabolizers.
  3. If the patient was previously on warfarin, start with home dose (or 10%-20% above home dose if INR less than 1.5).
  - Loading Dose Days 2-7
    1. Check INR in the morning; use each result to plan that evening’s warfarin dose:
      In the patient with baseline normal INR/PT, normal liver functions and who is not on medication interacting with warfarin, may check INR on days 3, 4, 5 and 7 or at clinician’s discretion
2. A reasonable method of adjusting warfarin doses according to INR is as follows:

   If INR is 1.1-1.3, repeat the initial loading dose*
   If INR is 1.4-1.9, reduce the initial loading dose by 50%
   If INR is 2-3, reduce the initial loading dose by 50%
   If INR is 3-4, reduce the initial loading dose by 25%
   If INR is 4-4.4, hold dose, check INR following day then resume warfarin at 50% of previous dose
   If INR >4.5, hold until INR <4.5 then resume warfarin at 50% of previous dose
   If INR >5.5, hold and check INR daily, when INR <5, restart at 25% less than previous dose or at discretion of primary clinician.

3. Experienced clinicians sometimes increase the designated initial loading dose 25-50% if the response to the chosen loading dose is very slow (e.g., little or no response after two doses). This maneuver shortens the time to therapeutic anticoagulation but increases the risk of a supratherapeutic INR.

4. In the patient whose baseline physiology and metabolism are stable, the dose on Day 8 of therapy can be calculated by dividing the total dose over the first week by 7, and one can proceed to recommendations for Long-Term Monitoring.

   o Once a patient has completed the initial phase, dosing is calculated over 7 days (i.e. total WEEKLY dose). Adjustments are made by increasing or decreasing the weekly dose between 5-20%.

   o Long Term Monitoring
      1. Routinely monitoring of INR once every 2-4 weeks in patients with:
         a. Stable INR
         b. No clinical indications for more frequent monitoring e.g:
            i. Started antibiotics or antiarrhythmics
            ii. Started on new medication known to interact with warfarin
            iii. Symptoms of bleeding
            iv. Significant changes in diet
            v. illness – particularly GI illness

   o Guidelines for adjustment of warfarin dose if INR outside of anticoagulation goal
      1. For maintaining INRs between 2.0 and 3.0
         If INR is 1.1-1.4, Check for compliance, if compliant, increase maintenance dose by 20%
         If INR is 1.5-1.9, increase maintenance dose by 10%
         If INR is 2-3, no change
         If INR is 3.1-4, decrease dose by 10%
         If INR is 4.1-4.5, decrease dose by 20%
         If INR >4.5, hold one dose, then restart 20% lower the previous dose*
         If INR >5, hold and check INR daily, when INR <4.5, restart at 25% less than the previous dose or at discretion of the primary clinician. *
2. For maintaining INRs between 2.5 and 3.5
   If INR is 1.1-2, Check for compliance, if compliant, increase maintenance dose by 20%
   If INR is 2-2.5, increase maintenance dose by 10%
   If INR is 2.5-3.5, no change
   If INR is 3.6-4.5, decrease dose by 10%
   If INR is 4.5-5, decrease dose by 20%
   If INR >5, hold one dose, then restart 20% lower than previous dose*

*Depending on the reason for anticoagulation and the perceived greater risk (risk of bleed versus risk of thrombus) dose reduction may be the preferred method of management over holding a dose to avoid a possible sub therapeutic level/ thrombus.

3. Additional guidelines
   a. When the INR is above the therapeutic range but <5, and the patient is asymptomatic (no clinically significant bleeding) and reversal is not required for surgical intervention, the dose of warfarin can be reduced or the next dose omitted and resumed (at a lower dose) when the INR approaches the desired range.

   b. If the INR is between 5 and 9 and the patient is not bleeding and has no risk factors that predispose to bleeding, the next 1 or 2 doses of warfarin can be omitted and warfarin reinstated at a lower dose when the INR falls into the therapeutic range. Alternatively, the next dose of warfarin may be omitted and vitamin K (1 to 2.5 mg) given orally. If vitamin K is given an INR should be checked at 12 hours after administration. This approach should be used if the patient is at increased risk of bleeding and their risk of a major adverse event from clotting is low.

   c. When more rapid reversal is required to allow urgent surgery or dental extraction, vitamin K can be given orally in a dose of 2 to 5 mg, anticipating reduction of the INR within 24 hours. An additional dose of 1 or 2 mg vitamin K can be given if the INR remains high after 24 hours.

   d. If the INR is 9 but clinically significant bleeding has not occurred, vitamin K1, 3 to 5 mg, should be given orally, anticipating that the INR will fall within 24 to 48 hours. The INR should be monitored closely and vitamin K repeated as necessary.

   e. When rapid reversal of anticoagulation is required because of serious bleeding or major warfarin overdose (e.g., INR 20), vitamin K1 should be given by slow intravenous infusion in a dose of 10 mg supplemented with transfusion of fresh plasma or prothrombin complex concentrate, according to the urgency of the situation. It may be necessary to give additional doses of vitamin K1 every 12 hours.
f. In cases of life-threatening bleeding or serious warfarin overdose, prothrombin complex concentrate replacement therapy is indicated, supplemented with 10 mg of vitamin K1 by slow intravenous infusion; this can be repeated, according to the INR. If warfarin is to be resumed after administration of high doses of vitamin K, then heparin can be given until the effects of vitamin K have been reversed and the patient again becomes responsive to warfarin.

**Associated Complications:**
- Risk of bleeding
- Teratogenic effects on fetus if mother is on warfarin during first trimester of pregnancy

**Special Considerations:**
- Subtherapeutic INRs: Inquire about missed doses or recent dose changes of existing medications, diet and exercise changes (increased exercise, or “healthy” dietary habits can lower an INR), new medications, herbal supplements or green tea consumption. Anecdotal resources indicate that marijuana use will lower INR (Potential Interactions - Alternative Therapies and Warfarin: Potential Interactions of Herbs with Warfarin [http://www.medscape.com/viewarticle/406896_2]

- Supratherapeutic INRs: Inquire about missed doses that have been caught up (i.e. two doses in <24 hour period), new medications or recent dose changes of existing medications, illness that affect appetite (sore throat, nausea/vomiting etc) and alcohol consumption.

- INR values are affected with many different drugs; antibiotics are the largest group of medications that will raise an INR. Managing practitioners should have INR checked 2-3 days after beginning a course of therapy to evaluate for elevation and potential dose adjustment and a repeat INR towards the end of the course for patients with an INR increase.

- Patients who have a low intake of vitamin K have been found to have more fluctuation in their INR. A consistent moderate intake or daily supplementation has been shown to decrease fluctuations in INR levels.

- Patients should avoid certain activities that are unsafe with taking an anticoagulant medication:
  - ATV riding
  - Boxing
  - Diving
  - Field Hockey
  - Football (tackle)
  - Ice Hockey
  - Motorcycle riding
  - Rock climbing
  - Rugby
  - Tae Kwon Do
  - Wrestling
Resources:

Children’s Hospital Boston Formulary, Boston, Massachusetts


Micromedex™ available online by subscription


Patient Resources:
http://www.clotcare.com/clotcare/ptinr.aspx (about the INR test)


http://www.stoptheclot.org/natt_publications/KandWarfarin.pdf (vitamin K and warfarin patient information)

http://www.clotcare.com/vitaminkandwarfarin.aspx (vitamin K and warfarin patient information)

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