A case of swollen hands

Patient Presentation
A 25 year old female presented to her primary healthcare physician complaining of swollen hands for the last week. She had also noticed that over the last two days she was getting swelling around her eyes and was short of breath.

History
History revealed that three weeks ago she had infected skin lesions on her thighs that had been treated with an antibiotic ointment from her doctor.

Past medical history
No past medical history of note. Recent HIV test was negative (2 months ago) with no history of unprotected sex since that time.

Past surgical history
No past surgical history.

Family History
Mother has hypertension and is on treatment
Father has no known medical concerns.
Four siblings, all healthy

**Allergies**

Non known

**Medication**

Multivitamins

**Travel History**

None noted

**Social history**

Not married, 1 child
Non smoker
No alcohol consumption
No illicit drug use

**Differential Diagnosis**

- Acute renal failure
- Congestive heart failure
- Immunoglobulin A (IgA) nephritis
- Membranoproliferative glomerulonephritis (MPGN)
- Lupus nephritis
- Post streptococcal glomerulonephritis (PSGN)

**Examination**

On Admission:

**Appearance**

Awake and alert, periorbital oedema, bilateral oedema of the hands and feet and mild tachypnoea at rest.

**Vitals**

Temperature: 37°C, afebrile
BP: 160/100 mmHg, hypertensive
HR: 85 beats per minute, tachycardic
RR: 20 breaths per minute, tachypnoeic

General
Mild conjunctival pallor
Peri orbital oedema and bilateral oedema of the hands and feet

Cardiovascular
Tachycardia
Hypertensive
JVP elevated to the angle of the jaw

Respiratory system
Mild tachypnoea
Bilateral basal crackles

Abdomen
Not distended
No tenderness
No organomegaly
Bowel sounds present

CNS
Normal level of consciousness, alert and co-operative
Gait, power, tone, sensation and reflexes intact and functioning within normal limits.

Dermatological
Mild scarring from healed lesions noted on thighs.

<table>
<thead>
<tr>
<th>Investigations</th>
<th>Results</th>
<th>Reference ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>WCC</td>
<td>5.31</td>
<td>4-12x10^9/L</td>
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<tr>
<td>HB</td>
<td>11.5</td>
<td>12.1-15.2g/L</td>
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<tr>
<td>Platelets</td>
<td>219</td>
<td>140-450x10^9/L</td>
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<tr>
<td>Normocytic normochromic anaemia</td>
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<tr>
<td>Urea</td>
<td>38</td>
<td>(2.5-6.4 mmol/L)</td>
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<tr>
<td>Creatinine</td>
<td>780</td>
<td>(50-80 umol/L)</td>
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<tr>
<td>Investigations</td>
<td>Results</td>
<td>Reference ranges</td>
</tr>
<tr>
<td>------------------------</td>
<td>---------</td>
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</tr>
<tr>
<td>K</td>
<td>6.1</td>
<td>(3.3-5.0 mmol/L)</td>
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<tr>
<td>Inorganic phosphate</td>
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<td>(0.8-1.4 mmol/l)</td>
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<tr>
<td>Calcium</td>
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<td>(2.1-2.6 mmol/l)</td>
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<tr>
<td>Complement</td>
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<td></td>
</tr>
<tr>
<td>CH50</td>
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<td></td>
</tr>
<tr>
<td>C3</td>
<td>Low</td>
<td></td>
</tr>
<tr>
<td>C4</td>
<td>Normal</td>
<td></td>
</tr>
<tr>
<td>ASOT</td>
<td>1280</td>
<td>(&gt;200 significant)</td>
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<tr>
<td>Anti-DNAse B</td>
<td>Positive to a titre of 360 (&gt;800 significant)</td>
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<tr>
<td>Blood Gas</td>
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<tr>
<td>pH</td>
<td>7.2</td>
<td>(7.38-7.42)</td>
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<tr>
<td>pCO2</td>
<td>3.8</td>
<td>(4.5-6.1 kPa)</td>
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<tr>
<td>Base excess</td>
<td>-17</td>
<td>(-2 to +2)</td>
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<tr>
<td>Bicarbonate</td>
<td>12</td>
<td>(22-30 mmol/l)</td>
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<tr>
<td>Anion gap</td>
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<td>(10-15)</td>
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<tr>
<td>Urine Dipstix</td>
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<td></td>
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<tr>
<td>Blood</td>
<td>4+</td>
<td>Negative</td>
</tr>
<tr>
<td>Protein</td>
<td>2+</td>
<td>Negative</td>
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<tr>
<td>Dysmorphic red cells</td>
<td>Present</td>
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</tbody>
</table>

**Urine:**

Oliguric, showing blood and protein on examination (see results table)

**Chest X ray:**

Fluid overload

**ECG**

Normal

**Further clinical course:**

The patient persisted with rapidly progressive renal failure, a glomerular filtration rate failing to normalize, persistent hypocomplementemia, haematuria and proteinuria.
A renal biopsy was ordered, with the following findings:

**Light microscopy**
- Hypercellularity of the glomeruli with enlarged glomerular tufts
- Neutrophils and monocytes
- Crescent formation noted in 30% of the glomeruli

**Immunofluorescence**
- Deposits of immunoglobulin G and C3 present along the glomerular capillary wall and mesangium.

**Electron microscopy**
- Occasional patchy thickening in the glomerular basement membrane
- Humps of electron-dense immune-type deposits noted in the glomerular basement membrane overlying the mesangium.

Consistent with poststreptococcal glomerulonephritis (PSGN)

**Discussion**

**Background**

The most common form of acute glomerulonephritis is a diffuse proliferative glomerulonephritis that follows infections caused by group A beta-haemolytic streptococci. A classic case follows a streptococcal infection by 7-12 days. This is the time needed for the development of antibodies. Our case study focuses on this disease progression and series of graphics explains how the inflammatory response is caused by an immune reaction following the entrapment of immune complexes in the glomerular capillary membrane. The resulting proliferation of the glomerular capillary endothelial cells and the mesangial cells causing swelling of the capillary membrane and the accompanying signs and symptoms of post streptococcal glomerulonephritis.

Before we discuss the immune response in relation to disease progression we will first look at the pathology involved in this disease and the healthy functioning kidney.

**Bacterial infection with Group A Streptococci**
Group A Streptococcus bacteria, most commonly *S. pyogenes*, typically cause superficial infections of the throat (pharyngitis) or skin (impetigo). Bacteraemia is usually resolved within a few days, however, renal complications may develop a week or two later as a result of the humoral immune response to blood borne bacterial proteins that have specifically become deposited or “planted” in the kidneys (nephritogenic antigens). These nephritogenic antigens are thought to mediate damage to the glomerular basement membrane which disrupts the filtering function and allows antigens to cross into the Bowman’s space.

The innate and adaptive immune response to *Streptococcal* infection

Innate immune cells such as macrophages and neutrophils migrate to sites of infection and help to destroy bacteria. The adaptive immune system is also activated with the recruitment of CD4+ helper T cells and B lymphocytes responding to bacterial antigens. An important factor in the development of kidney dysfunction following a *Streptococcal* infection is the generation of antibodies that target the secreted bacterial antigens.
There are two possible explanations of how immune complexes become deposited in the kidney.

As plasma cells produce more and more antibodies, antibody-antigen complexes can form in the bloodstream due to the presence of secreted antigens and antibodies. These immune complexes can then become deposited in the kidneys via the circulation. Alternatively antibodies bind to free antigen already trapped in the glomerular basement membrane to form “in situ” immune complexes during glomerular filtration due to disrupted barrier function between the capillaries and the Bowman’s capsule.

To better understand the complications caused by immune complex deposition in the kidneys we will briefly look at the structure and functioning of a healthy kidney.

The normal filtering of waste products from the blood is performed by thousands of nephrons located in the kidney. These are composed of several structures including renal corpuscles, responsible for filtering large amounts of fluid from the blood.
The renal corpuscle consists of two main parts: the glomerulus and Bowman’s capsule. The glomerulus is a fine network of capillaries that are permeable to small molecules. The glomerular capillary membrane is similar to that of other capillaries except that it has three major layers instead of the usual two. The (1) endothelium, (2) basement membrane and (3) layer of epithelial cells (continuous with the epithelium lining the Bowman’s capsule) known as podocytes, that surround the outer surface of the capillary basement membrane. Podocytes have long processes (foot processes) that wrap themselves around the glomerular capillaries. Each component brings its own unique function- the endothelium is fenestrated, the basement membrane has a negative electrical charge and prevents the passage of plasma proteins and the podocyte foot processes form slit-pores – that together form a highly efficient filtration system. Fluid from the glomerular capillaries leaks out across these layers into Bowman’s capsule that surrounds the glomerulus. This glomerular filtrate contains only water and small molecules. Another important component of the renal corpuscle is the mesangium. This occurs in areas where the capillary endothelium and basement membrane do not completely surround each capillary. The mesangial cells instead provide support in these areas by producing an intercellular substance similar to that of the basement membrane. They also possess phagocytic properties and remove materials that enter the intercapillary spaces. Typically this area is narrow and contains only a small number of cells, however mesangial hyperplasia and increased mesangial matrix occur in a number of glomerular diseases.

Now that we have an understanding of the anatomy of the renal corpuscle we can look more closely at how streptococcal antigen is deposited in the kidney.
Two “nephritogenic” proteins have thus far been identified in Streptococcal infections and include SpeB, a bacterial serine protease enzyme, and NAPIr, a secreted bacterial protein known as “nephritis-associated plasmin receptor”. It is thought that these proteins when present in the kidney precipitate enzymatic damage to the basement membrane and endothelial cell integrity, thus allowing plasma proteins, erythrocytes and immune complexes to cross into the bowman’s space i.e. the system becomes “leaky”. The bacterial antigens bind and activate plasmin or plasminogen which in turn activates the enzymes collagenase and matrix metalloproteinases that degrade the basement membrane and weaken endothelial cell tight-junctions. Disrupted membrane function allows bacterial antigens to be deposited inside the Bowman’s space.

Immune complexes can be formed inside the Bowman’s space when free IgG antibodies cross the disrupted basement membrane and contact “planted” bacterial antigens already present, or alternatively, immune complexes that have formed in the bloodstream can also cross the weakened barrier and become trapped in the Bowman’s space.
**Complement activation in the kidney**

The deposition of immune complexes in the Bowman’s space activates the classical complement system by recruitment of C1 proteins from plasma. A loss of plasma C3 levels (see investigations) correlates with complement activation and deposits of C3 proteins can be detected along the basement membrane and in the Bowman’s space.

**Immune cell recruitment to the kidney**

Due to complement activation and generation of anaphylatoxins we then see an increased influx of immune cells such as macrophages and neutrophils. Deposits of C3, antibody and bacterial antigens accumulate at the basement membrane near the podocyte foot processes, known as “humps”. Kidney function is severely hampered by influx of immune cells, particularly neutrophils, that can block capillaries. There is also marked cell proliferation of mesangial and endothelial cells due to the presence of soluble inflammatory molecules.
Kidney function is compromised by the breakdown of the filtering function of the basement membrane and the influx of numerous immune cells due to complement activation. This results in plasma protein (proteinuria) and erythrocyte (haematuria) leakage unto the urine and a decrease in urine volume (oliguria). Oliguria which develops as the glomerular filtration rate (GFR) decreases is one of the first symptoms. Proteinuria and haematuria follow because of increased glomerular capillary wall permeability. The red blood cells are degraded and cola coloured urine is one of the early signs of the disorder. Sodium and water retention gives rise to oedema, particularly in the face and hands and the patient is also found to have hypertension due to salt imbalance. Laboratory findings include elevated antistreptolysin O titre (antibodies to a streptococcal exoenzyme) a decline in C3 complement and cryoglobulins. In most cases only mild damage to the kidneys is sustained and the patient will recover, but ongoing inflammation can result in complete and irreversible kidney damage that requires chronic dialysis and ultimately a transplant of a functional kidney.

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**Treatment**

Symptomatic therapy is recommended for patients with acute poststreptococcal glomerulonephritis, and it should be based on the clinical severity of the illness. The major goal is to control fluid balance and blood pressure.

This patient did not respond to diuretics, remaining fluid overloaded with pulmonary oedema and persistent oliguria. The patient required urgent dialysis, which was continued for three weeks.
Final Outcome

The patient did not recover from her three weeks of dialysis.

She was then assessed for the chronic dialysis program and work up was started for the transplant program.

References


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Evaluation - Questions & answers

What was the final diagnosis?
Post streptococcal glomerulonephritis (PSGN)

What is an important factor in the development of kidney dysfunction following a streptococcal infection?
The generation of an abundance of antibodies that target the secreted bacterial antigens, forming immune complexes that then deposit in the renal corpuscle of the kidney.

What are the three layers that make up the filtration membrane of the glomerulus?
The endothelium, basement membrane and layer of epithelial cells (continuous with the epithelium lining the Bowman’s capsule) known as podocytes.

How are the two nephritogenic proteins SpeB and NAPIr thought to contribute to PSGN?
When these proteins are present in the kidney they precipitate enzymatic damage to the basement membrane and endothelial cell integrity, thus allowing plasma proteins, antibodies, erythrocytes and immune complexes to cross into the Bowman’s space. Disrupted membrane function allows bacterial antigens to be deposited inside the Bowman’s space where antibodies can detect them and generate “in situ” immune complexes.

How does the basement membrane become degraded?
The bacterial antigens bind and activate plasmin or plasminogen which in turn activates the enzymes collagenase and matrix metalloproteinases that degrade the basement membrane and weaken tight junctions between endothelial cells.

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Multiple Choice Questions

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