

A pepper-pot skull?

HOSP #		WARD	General Practitioner Practice in Robertson
CONSULTANT	Dr. Jody Rusch	DOB/AGE	83 year Male

Abnormal Result

Serum protein electrophoresis demonstrates a 4.4 g/L, IgG kappa monoclonal peak in the gamma region.

Presenting Complaint

Complains of bilateral hip pain and RUQ discomfort.

History

Atrial fibrillation on Xarelto.

2 x CABG

Examination

RUQ pain and tenderness

Hear rate regular

Laboratory Investigations

Urine protein electrophoresis: No Bence Jones protein

Serum free light chains:•

- Kappa 62.87 mg/L (3.30-19.40)•
- Lambda 19.63 (5.71-26.30)•

- K:L ratio 3.20 (0.26-1.65)
- Creatinine 108 (eGFR 56)
- Calcium 2.42 mmol/l
- Albumin 40 g/L
- Hb 12.7 (11.0-16.0)

Other Investigations

U/S shows gallstones.

X-Ray of pelvis shows “sclerotic changes to both hips and pelvis”

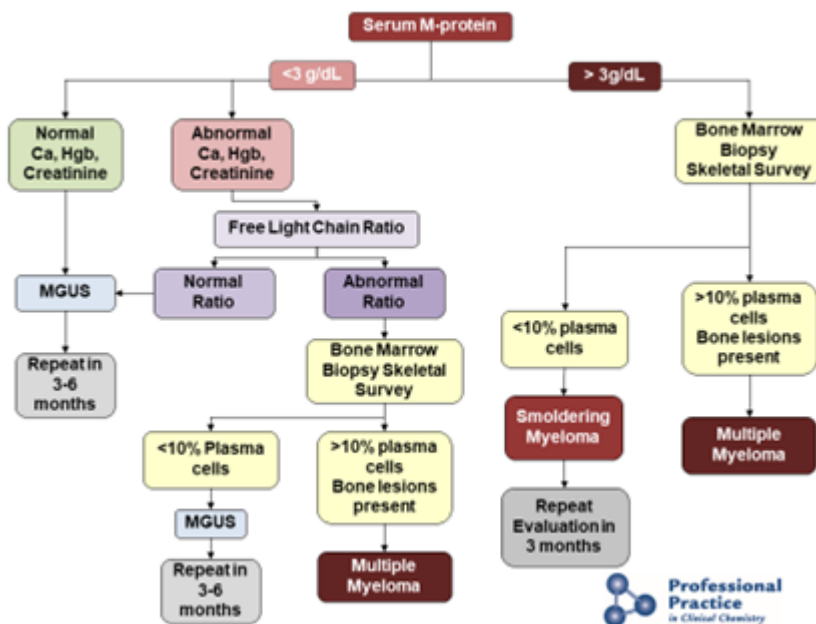
Final Diagnosis and Take Home Message:

1. What is the likely diagnosis

This 83 year old male with multiple co-morbidities presenting with signs and symptoms suggestive of multiple myeloma, confirmed on SPE as IgG Kappa.

- CRAB criteria before performing SPE: C- R+ A- B+ (2/4)
- Bony pain could be secondary to lytic bone lesions associated with MM, but also possibly to due sclerotic/wear-and-tear when considering his age. RUQ pain is likely due to gallstones.
- Renal impairment – this is probably normal renal function for an 83 year old man
- In medicine generally an eGFR < 60 is representing renal impairment (stage 3)
- However, in monoclonal disease eGFR < 40 or serum Cr > 177 is the cut-point
- Bone lesions – myeloma classically causes lytic bone lesions, e.g. “pepper-pot skull”

It was suggested that the clinician talks to the radiologist as to whether the X-Rays were in keeping with myeloma.



2. Critically discuss whether this patient needs a bone marrow biopsy.

The patient's age along with co-morbidities would concern any drastic intervention:

- he will be an anesthetic risk for BM Bx to be performed in theatre (assuming that is standard procedure), and
- will the BM biopsy give add anything further to the already established IgG Kappa diagnosis, which can be treated accordingly.

The case should ideally be discussed with Oncology. A bone marrow biopsy is done under local anaesthetic. The bone marrow will allow the haematologist / oncologist to assess the degree of marrow clonal infiltration. The important cut-offs are 10 & 60%. This is important to decide on diagnosis, stage, prognosis, treatment and later, the response to treatment. The criteria for doing a bone marrow biopsy at our centre are:

- Positive CRAB.
- IgG monoclonal peak > 15 g/L.
- IgM or IgA monoclonal.
- FLC K:L > 10

Why is there a lower (10%) limit for degree of marrow clonal infiltration? Is there a link to immunoparesis? One likely always has some clonal expansion in bone marrow, probably a normal or a non-pathological finding.

3. Discuss the serum FLC in the setting of the renal impairment.

FLC are filtered and reabsorbed by the nephron under normal circumstances, along with other LMW proteins. During a plasma cell dyscrasia, the nephron is overwhelmed by the amount of FLC (stemming from monoclonal origin), can cause renal impairment. Hence, renal function being part of the CRAB criteria. Furthermore, renal impairment itself (in the absence of MM), can cause elevated Kappa and Lambda FLC – usually with a slight higher ratio =3.2.

In patients with renal failure, there is greater retention of serum free light chains. It is difficult to interpret ratios ranging between 1.65 – 3.0 in the context of renal insufficiency. In such cases, further investigation with a 24-hour urine protein electrophoresis and urine immunofixation helps to guide interpretation. If both of these subsequent studies are normal and the patient has no other symptoms suggestive of a plasma cell dyscrasia, then the increased ratio is likely due to the renal insufficiency.

4. Discuss electrophoresis briefly.

Electrophoresis is a general term that describes the separation of charged particles/ ions under the influence of

an electric field – in this case the charge of proteins. Migration of proteins is based on their charge, size and velocity (product of their mobility and field strength) Make sure you understand why the proteins are charged the importance of NET charge and how we keep those charges stable in the field. If I can take a crack at this: The overall NET charge of the molecule is based on the number of elements (incl. amino acids with varying side-chains moieties) (I think this is the confusion when some mention that electrophoresis is based on charge, and also size. I don't necessarily think that the two are synonymous), and each amino acid has different degrees of charge based on their differing R-group. The stability of the charges within the field is achieved by running the sample solution through a buffer. Right about the buffer. Remember that size and charge are two different physical aspects that you can use to separate molecules. For example, a DNA gel is a separation purely based on size. The net charge is the same on all the molecules. The net charge in proteins is from the side chains, which is why you have to learn about neutral, acidic and basic amino acids. The side chains have different pKa's and so are charged differently.

a. What is the difference between capillary and gel electrophoresis. Explain which your lab uses and why.

What I described in Q4 was basically the concept of gel electrophoresis where agarose gel is used as the medium in which the proteins are separated according to their size, charge, and interaction with the medium itself. At TBH we use gel electrophoresis, but will soon be getting a Minicap/ CZE. CZE: As with gel electrophoresis, CZE also separates ions based on their electrophoretic mobility with the use of an applied voltage – all dependent on the charge of the molecule, viscosity and particle size. CZE's voltage is much higher compared to GE – quicker results. The buffer/ mobile phase of

CZE uses an electrolyte filled capillary, where eletro-osmotic flow (EOF) is generated: similarly sized and charged ions move together and are subsequently separated and detected at different time intervals. The more voltage you apply the faster the separation occurs. However, the limiting factor is that applying high voltages generates a lot of heat which can denature proteins, thereby altering their conformational shape and changing their NET charge. Capillaries are much more effective at shedding heat as they are long and thin. Thus, very high voltages can be applied and the run time is much shorter. In gel electrophoresis, you measure how far the molecules travel in a set time, e.g. 1 hour. In capillary electrophoresis, the distance that the molecules move is set and so you measure the time it takes for the molecules to travel that set distance (like running a 100m race).

The way I reconcile how the CZE differential seperation works is by the

1. driving force of the buffer through the tubing (forward force)
2. negative charge on the side of the tube (retarding force)
3. NET charge on the molecule (many amino acids=higher charge, eg Albumin) (determine degree of retardation of flow)
4. Voltage powers the whole system

5. Why is the serum FLC abnormal but not the urine protein electrophoresis?

UPE's sensitivity is limited due to the reabsorption of FLC in the renal tubules. FLC in urine will only be detected until loss of tubular function/ tubules are overwhelmed by FLC volumes. This patient's Kappa FLC of 63 mg/L in serum should be detected on UPE, but tubular function is seemingly still

intact with little being excreted.

Some are of the opinion that SPE and SFLC is the preferred method to screen for myeloma because of higher sensitivity and specificity, as opposed to SPE and UPE, which may have a slightly lower sensitivity.

It should however be noted that quoted sensitivities and specificities are usually based on retrospective audits of patients who eventually end up in a myeloma clinic. So, it is not sure what the sensitivity and specificity is if you just screen the general population, older people, people with some vague symptoms...

6. Against which epitopes are the FLC assay directed?

The FLC epitopes are located between the interface between the light and heavy chains and are “hidden” – when bound to Ig, they will not be detected. Only when these epitopes are “free”, can they be detected, hence free light chains. They are directed at 2 hidden epitopes.

7. Why is the FLC assay polyclonal and not monoclonal?

The biggest decider many times is COST, but lets put that aside for now.

It appears that polyclonal assays are more robust and have higher yields in product during testing and easier to make. They are unfortunately less specific, but this is not the most critical when one wants to measure the FLC broadly, instead of particularly specific sites.

Epitopes are three dimensional shapes that the antibody binds to. This is determined by the amino acid sequence. One drawback with polyclonal assays is that lot to lot will vary.

The difficulty is to maintain consistency in further production and / or distribution of the antibody – it is not a simple process to ensure consistency.

8. Describe how a monoclonal antibody is made for use in an assay.

Inject a rabbit (or other animal) with the protein of choice. In three weeks, the rabbit will have produced antibodies to the protein. The rabbit is sacrificed (killed) and the spleen harvested. The spleen is ground up and the cells are put in a culture with a certain myeloma cell line. The culture medium contains colchicine that induces the rabbit cells and the myeloma cells to fuse. It also contains HAT medium: hypoxanthine, aminopterin and thymidine. This specific myeloma cell line cannot recycle thymidine in the presence of hypoxanthine.

So in the culture there are now three cell lines.

- Firstly, there are the rabbit cells that haven't fused; these will die because they are not immortal.
- Secondly, there is the myeloma cell line, this will also die because of the recycling problem.
- Finally, there is a fused cell line that will survive .

Each of these surviving cell lines will produce one Ig against one part of the protein. Now the researchers take the medium and put a tiny amount into a well. The amount is so small that *on average* each well will contain only one cell; some will of course contain nothing. Then, each well is targeted against the protein and the most promising ones are investigated further. An immortal Ig producing factory directed against one epitope and based on one cell line, a single clone, or as we'd call it a monoclonal, has been produced. Each manufacturer's produced immunoglobulin is different and may produce better or worse results.

9. The GP, in Robertson, wants some advice on how to proceed. What do you tell him?

A multidisciplinary approach would be best:

- Treatment for the lytic bone lesions (after opinion by radiology): bisphosphonate
- Assess overall medication and lifestyle to determine overall risk for worsening renal dysfunction (drugs, co-morbidities.....always suggest stopping smoking/drinking)
- Prevention of thrombotic/infective episodes
- Treatment of any further abnormalities should they arise (hypercalcaemia, anaemia etc.)
- Specialist referral:
 - Haematoncologist for treatment of MM: UPE , Bone marrow biopsy
 - General surgery for gallstone

10. Is there any relevance for the RUQ/gallstone pain in myeloma specifically?

There are some reports where cholecystitis has presented in MM (mets etc), but it is not a separate entity on its own (such as in POEMS), this is simply the real world where elderly patients have more than one pathology.