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How Measuring Glomerular Filtration Rate? Comparison of Reference Methods

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1. Introduction

Glomerular filtration rate (GFR) is considered as the best way to assess global renal function (Gaspari et al., 1997; Stevens & Levey, 2009). Even if GFR estimations (based on creatinine- or cystatin C-based equations) are most often used (see Table 1)(Cockcroft & Gault, 1976; Levey et al., 1999; Levey et al., 2006; Levey et al., 2009), measuring "true" GFR is still important in clinical practice, especially in particular patients (Delanaye et al., 2011a; Delanaye & Cohen, 2008; Stevens & Levey, 2009). In this chapter, we will review the different markers which can be considered as reference methods to measure GFR. Before moving to clinical trials, we have to recall the physiological characteristics of an ideal GFR marker.

2. Clearance concept and ideal marker for glomerular filtration rate

The history of the renal physiology is deeply influenced by the book published by Homer W. Smith in 1951 (Figure 1): «The kidney: structure and function in health and disease »(Smith, 1951b). In this best-seller of nephrology, Smith compiled all the physiological data (more than 2300 references) which have been published in the scientific literature until 1951. Smith, himself, has largely contributed to the physiological knowledge of the kidney. A large part of this book is dedicated to the GFR measurement. The concept of clearance is well explicated. Actually, the Danish physiologist, Poul Brandt Rheberg was the first to use and define the concept of clearance in 1926 even if this author did not use the word "clearance". Rheberg studied on himself the urea and creatinine clearances to prove that kidney has a filtrating and not only a secreting action (Rehberg, 1926b; Rehberg, 1926a). The term clearance was used for the first time by Möller in 1929 and was then concerning the urea clearance which was proposed as the first evaluation of renal function (Möller et al., 1929). Smith has largely contributed to make popular and classical this concept of clearance to assess GFR (Smith, 1951a). Renal clearance of a substance is defined as the volume of plasma cleared from this substance per time unit (mL/min). Clearance is thus a virtual volume but will permit to apprehend GFR and renal function. However, the concept of clearance is applicable to any internal or external substances. To be considered as a reference method, a marker must have strict physiological characteristics (Smith, 1951b):

1. Marker production and marker plasma concentration must be constant if GFR does not change

- 2. Marker must be free in plasma (not binding to protein) and must be freely and fully filtrated through the glomerulus
- 3. Marker is neither secreted nor absorbed by renal tubules
- 4. Marker must be inert and, of course, not toxic
- 5. Marker excretion must be exclusively excreted by kidneys
- 6. Marker must be easily measured in both plasma and urine

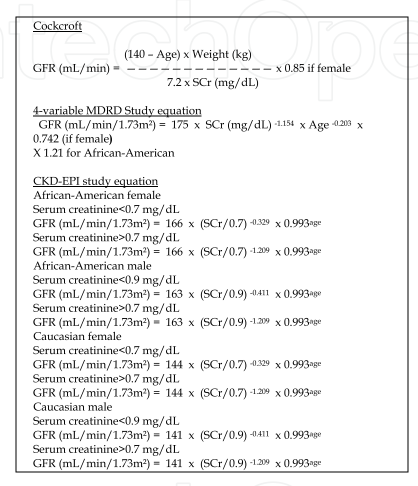


Table 1. Creatinine-based equations. SCr: Serum Creatinine, GFR: glomerular filtration rate, MDRD: Modified diet in renal disease, CKD-EPI: Chronic Kidney Disease-Epidemiology group.



Fig. 1. Homer W. Smith

The renal clearance will be easily calculated with the following equation:

$$GFR=([U] \times V) / [P]$$
 (where $[U]$ = urinary concentration, $[P]$ = plasma concentration, V = urinary volume)

The calculated value will be then divided by the time interval where the urine collection has been made. *Sensu strict*, the plasma concentration must be sampled from arterial blood but errors induced by venous samples are very limited (Laake, 1954; Handelsman & Sass, 1956; Nosslin, 1965). In the same view, the transit time through the urinary system should also be taken into consideration but, once again, error linked to this transit time is negligible (Ladegaard-Pedersen, 1972; Nosslin, 1965). The method originally proposed by Smith for measuring GFR is not an easy task. Actually, the marker (inulin see below) must be intravenously injected and then perfused at a constant rate to reach stable plasma concentrations. Thereafter, urine collection must be realized, which is a potential source of errors. For this reason, Smith recommended urine collection on 10 and 15 minutes with the use of urinary catheter. Smith recommended three successive collections. The patient was hydrated to assume a sufficient urinary flow though these collections. The mean of the three collection was considered as the GFR measurement (Smith, 1951a). Nowadays, the urine collections are done without urinary catheter and on a longer period of time (60 minutes) to decrease the impact of urine collection errors on the final result (Levey et al., 1991; Robson et al., 1949).

The ideal marker does not exist in the organism (or has still not been discovered if we want to be optimistic). Both urea and creatinine clearance have strong limitations, notably because creatinine is secreted and urea is absorbed by renal tubules (Dodge et al., 1967; Morgan et al., 1978). Therefore, exogenous markers are used to measure GFR. We will successively describe the markers which are still used in clinical practice in 2011: inulin, ⁵¹Cr-EDTA, ⁹⁹Tc-DTPA, iothalamate and iohexol. For every marker, we will describe strengths and limitations both from an analytical and clinical point of view.

3. Inulin

Inulin is still considered nowadays as the gold standard to measure GFR. Smith has deeply studied this marker and makes it the most popular. Inulin is a polymer of fructose which is found in some plants which uses it as energy provider in place of amidon. Its molecular weight is 5200 Da (Gaspari et al., 1997). Some plants are especially rich in inulin: chicory, garlic, leek and Jerusalem artichoke. Humans are not able to metabolize inulin. Because inulin is the first reference method to have been used, its role in the GFR measurement has only be asserted on basis of physiological studies (because the first method is not comparable to any other!). Once again, we often refer to the studies published by Smith and Shannon (New York university)(Smith, 1951a; Smith, 1951c) and by another pioneer Richards (Philadelphia university)(Richards et al., 1934). Inulin was obviously considered as a safe product with any effect on GFR (Shannon, 1934). Inulin is freely filtrated through a semi-permeable membrane which is a strong argument for the absence of binding to protein. This has been shown by Shannon in 1934 (Shannon, 1934) and by Richards in 1937 (Hendrix et al., 1937). In the same publication, Richards proved that inulin was freely and fully filtrated through the glomerulus because he measured the same inulin concentration both in the plasma and the glomerulus of a frog and a salamander (Hendrix et al., 1937). The absence of both tubular absorption and secretion has been demonstrated by an important article published by Shannon in 1934 (Shannon, 1934). In this article, this author showed the

absence of inulin excretion in two types of aglomerular fishes (goosefish, Lophius piscatorius and toadfish Osteichthyes - Lophiidae). In the same article, Shannon measured GFR by inulin clearance in another type of fish with glomerulus, the dogfish (Chondrichthyes - Squalidae). These fishes were then treated with phlorizin which was sensed to block all tubular activity. Although the creatinine clearance in this fish was increased, the inulin clearance was not modified by this treatment (Shannon, 1934). In the same year of 1934, inulin clearance was also measured in aglomerular fish and in dogs by Richards (Richards et al., 1934). The experimentation (measuring GFR with and without phlorizin) was then repeated in man by Smith and Shannon. The results obtained in animals were confirmed in humans. Shannon was the first human who was perfused by inulin in 1935 (Shannon & Smith, 1935; Smith, 1951c). These authors had thus suggested that inulin was not secreted by renal tubules. This assertion will be thereafter confirmed by other authors with the same type of methodology (Shannon & Smith, 1935; Alving et al., 1939; Laake, 1954). Additional arguments were developed in the sixties by animal studies using micropontions in the tubules (Gutman et al., 1965). After intravenous injection, inulin is fully excreted by kidneys in urine (Shannon & Smith, 1935), even if very low concentrations of inulin are found in bile (Höber, 1930; Schanker & Hogben, 1961).

Inulin is doubtless the marker who has been the most investigated from a physiological point of view. In this view, it is logical that inulin is still considered as the gold standard for GFR measurement. Nevertheless, there are limitations to its use in daily practice. Because its relatively high molecular weight (5200 Da), the molecule is relatively viscous and don't quickly reach its volume of distribution. Therefore, only methods using urinary clearance with constant infusion rate seem accurate for this marker. Such methods are more cumbersome. Moreover, inulin is not easily available on the market and remains relatively costly. From our point of view, the most important limitation of inulin is the difficulty linked to its measurement in urine and plasma. Actually, several methods have been proposed and these methods are probably not interchangeable. There is no standardization in inulin measurement. We have shown that GFR results could vary from -10 to +10 mL/min in the same patient only because inulin was measured by a different method (unpublished data). Moreover, most of the methods (except the enzymatic ones) are prone to interferences with glucose measurement which is a limiting factor when measuring GFR in diabetic patients (Little, 1949). Regarding the methods for measuring inulin, we can cite the "acid" methods (Kuehnle et al., 1992; Shaffer & Somogoyi, 1933; Alving et al., 1939; Corcoran, 1952; Rolf et al., 1949; Roe, 1934; Steinitz, 1938; Hubbard & Loomis, 1942; Lentjes et al., 1994; Heyrovsky, 1956; Rolf et al., 1949), the enzymatic methods (Day & Workman, 1984; Delanghe et al., 1991; Jung et al., 1990; Summerfield et al., 1993; Dubourg et al., 2010) and the new methods by high performance liquid chromatography (HPLC) (Ruo et al., 1991; Baccard et al., 1999; Dall'Amico et al., 1995; Pastore et al., 2001). Describing these methods in detail are beyond the scope of this chapter and we propose the readers the following reference if they are interested in this topic (Delanaye et al., 2011b).

4. Preliminary statistical considerations

The use of inulin as GFR marker is justified by physiologic studies. The others markers that will be proposed thereafter will be compared to inulin measurements. Therefore, the use of other markers will be justified not by physiological studies (even if some

physiological studies exist for some markers) but by studies comparing these markers with inulin. Unhopefully, most of these studies comparing different GFR tests lack of strong statistical methodology. Actually, most of the authors have only shown a good correlation between the markers, which is expected but not sufficient. Ratio of new markers results on inulin results are also used (the result being considered as good if ratio is near to 1). The use of such ratio may be misleading (for example, if one method overestimates true GFR in low GFR levels but underestimates GFR in high levels, the ratio will be near to 1 although the method is actually not precise enough). To compare the performance of a new GFR measurement compared to inulin, we need to know the bias (mean difference between the two results) and the precision (standard deviation (SD) around the bias) of this new measurement. Bland and Altman analysis is thus required (Bland & Altman, 1986).

Regarding the other GFR markers, we must also stress that GFR can be measured by plasma clearance and using a bolus injection (instead of constant infusion rate) which makes the GFR measurement much more simple. Method to measure GFR by plasma clearances can be very different (number of samples, timing of samples, mathematical model used). We must keep in mind that results of plasma and urinary clearances are not strictly comparable (plasma clearances overestimate urinary clearances even if the overestimation decreases if plasma samples are drawn after 24 hours) and this must be integrated when these GFR methods are compared (Agarwal et al., 2009; Stolz et al., 2010).

5. ⁵¹Cr-EDTA (Ethylenediaminetetra-acetic acid)

5.1 Physiological and analytical data

⁵¹Cr-EDTA is an isotopic marker which has a low molecular weight (292 Da). Most of the authors consider that 51Cr-EDTA is not binding to proteins (<0,5% (Brochner-Mortensen, 1978; Bailey et al., 1970; Garnett et al., 1967; Stacy & Thorburn, 1966; Forland et al., 1966; Kempi & Persson, 1975; Forland et al., 1966)) even if Rehling described a binding to protein of 10% (Rehling et al., 1995; Rehling et al., 2001). Due to its low molecular weight, 51Cr-EDTA is freely filtrated through the glomerulus. Physiological studies about renal handling of 51Cr-EDTA are few but it seems that 51Cr-EDTA is neither secreted nor absorbed by renal tubules (Eide, 1970). This absence of secretion and absorption is also confirmed by Forland in dogs (Forland et al., 1966). Regarding the potential extra-renal excretion of 51Cr-EDTA, Garnett described a salivary and a fecal excretion under 1% in one anephric patient (Garnett et al., 1967). Brochner-Mortensen later confirmed the poor fecal excretion (less than 0.1% of the injected dose). Studying the renal excretion and the corporal global radioactivity of 8 healthy subjects after 72 hours, Brochner-Mortensen estimated that 4.5% of the 51Cr-EDTA will be retained in the body, especially in the liver and kidneys (Brochner-Mortensen et al., 1969). The difference between 51Cr-EDTA total clearance and 51Cr-EDTA urinary clearance corresponds to extra-renal clearance of the marker. With this methodology, the same authors estimated extra-renal clearance at 4 mL/min (and this extra-renal clearance remains stable for all GFR ranges)(Brochner-Mortensen & Rodbro, 1976). Jagenburg had also calculated an extra-renal clearance of 2 mL/min in two anuric dialysis patients (Jagenburg et al., 1978). Only, Rehling described a higher extra-renal clearance at 8.4% (Rehling et al., 1995).

Measurement of ⁵¹Cr-EDTA by nuclear count is very precise and easy because ⁵¹Cr-EDTA half time is long (27 days)(Chantler et al., 1969). The quantity of ⁵¹Cr-EDTA injected is

relatively small and therefore the irradiating dose received by the patient is very limited (absorbed dose from 0.011 to 0.0077 mSv according to the radioactive dose injected which is usually 7 MBq). This absorbed dose corresponds to the natural dose of irradiation received in one week and is much lesser than the dose received after thoracic radiography (0.02 mS). Nevertheless, we do not recommend this technique to measure GFR in pregnant women even if authors seem to use it safely (Brochner-Mortensen, 1978; Medeiros et al., 2009; Durand et al., 2006). The dose of EDTA is 1000x lesser than the dose considered as safe (Chantler et al., 1969).

5.2 Clinical data

The first studies about 51Cr-EDTA have been published in the sixties, even if studies (but with questionable methodology) had been published before with EDTA marked with ¹⁴Cr (Spencer et al., 1958; Foreman & Trujillo, 1954). In 1964, Downes was the first to give 51Cr-EDTA to cows to study the intestinal transit (Downes & Mcdonald, 1964). In 1966, Stacy and Thorburn are the first to inject 51Cr-EDTA to lambs for measuring GFR. They reported a good correlation with inulin clearance in the animal model (ratio 51Cr-EDTA/inulin was 0,95)(Stacy & Thorburn, 1966). The first scientists who will be interested in GFR measurement by 51Cr-EDTA in humans are English (Garnett et al., 1967; Favre & Wing, 1968; Garnett et al., 1967; Heath et al., 1968; Lavender et al., 1969). It must be underlined that nearly all studies published on this marker are coming from Europe because 51Cr-EDTA is not available in USA (not approved by the FDA)(Brandstrom et al., 1998). The first author who studied 51Cr-EDTA in humans is Garnett who was nuclearist in Southampton. These first data were published in The Lancet in 1967 (Garnett et al., 1967). This author injected one unique dose of 51Cr-EDTA and described a mono-exponential decrease in 51Cr-EDTA concentrations after 30 minutes. This author already evoked the plasma clearance (and the bolus injection) to measure GFR with 51Cr-EDTA. Unhopefully, Garnett did not compare his results to inulin clearance but only to creatinine clearance. However, Garnett performed and compared 56 51Cr-EDTA urinary clearances with inulin urinary clearances. He found a correlation of 0.995 and asserted that 51Cr-EDTA result were between ±5% of the inulin results which was really excellent. Thereafter, several studies were published on the same topic to compare performances of inulin clearance with urinary or plasma clearance of 51Cr-EDTA. We resumed these studies in Table 2, restricting the data to studies in adults. However, once again, the following conclusions will be drawn from studies having used the most adequate statistical methods. Globally, the performance of 51Cr-EDTA is good. Chantler, in 1969, showed that results of urinary clearance of 51Cr-EDTA was within 5% of the results of inulin (Chantler et al., 1969). This excellent concordance between urinary clearances of 51Cr-EDTA and inulin will be later confirmed by Froissart. This author showed a bias of +3 mL/min (51Cr-EDTA thus slightly overestimating inulin) and a precision of ± 4 mL/min (95% of the 51Cr-EDTA results will be + or - 8 mL/min around the bias)(Froissart et al., 2005b). The best study comparing 51Cr-EDTA plasma clearance with inulin clearance is certainly published by Medeiros in 2009 (Medeiros et al., 2009). This author showed that bias between the two GFR was 3±6 mL/min. This is one of the rare studies where accuracy 30% results are given (defined as the percentage of patients having a 51Cr-EDTA GFR within 30% of inulin GFR). Accuracy 30% for plasmatic clearance of 51Cr-EDTA is 93%. The higher performance is obtained when late blood samples (at 6 or 8 h) are considered.

References	Sample	Population	GFR range (mL/min/ 1.73 m²)	GFR methods	Statistics	Results
(Garnett et al., 1967)	56	NA	± 0 to 180	Urinary clearance and constant infused rate	Regression Correlation	=1.075x-3.06 0.995
(Heath et al., 1968)	39	Healthy CKD Calcium troubles	10 to 150	Urinary clearance and constant infused rate	Correlation	0.995 ⁵¹ Cr-EDTA underestimat es by de 14- 16%
(Favre & Wing, 1968)	20	CKD	6 to 187	Urinary clearance and constant infused rate	Ratio Correlation BAr	1.02 0.992 1.5±8.7
(Lavender et al., 1969)	100 clearances in 28 subjects	CKD	± 0 to 150	Urinary clearance and constant infused rate	Ratio Regression Correlation	0.96 ± 0.0027 =0.96x+0.26 0.994
(Brochner- Mortensen et al., 1969)	17	2 healthy	± 10 to 130	Inulin: urinary clearance and constant infused rate 51Cr-EDTA: plasmatic clearance: on 5 hours, samples every 15 min	Correlation Regression	0.974 =1.017x+1.6
(Chantler et al., 1969)	21	CKD	± 10 to 160	Urinary clearance and constant infused rate	Correlation Regression Ratio	0.977 =1.004x-0.032 1.004±0.013
(Stamp et al., 1970)	65 clearances in 56 subjects	15 healthy 41 calcium troubles	± 20 to 140	Urinary clearance and constant infused rate	Correlation Regression Ratio	0.91 =0.98x+6.5 0.96±0.02
(Ditzel et al., 1972)	20	NA	6 to 166	Inulin: urinary clearance and constant infused rate ⁵¹ Cr-EDTA: plasmatic clearance: samples at 5,10, 15, 20, 30, 60, 90, 120, 150, 180, 210, 240 min	Correlation Regression BAr	0.97 =0.85x+11.42 1.5±11.7
(Lingardh, 1972)	25	Healthy and CKD	±8 to 120	Inulin: urinary clearance and constant infused rate ⁵¹ Cr-EDTA: plasmatic clearance: samples timing not available	Correlation Regression Mean difference	0.984 =1.099x+4.96 6.2 mL/min

(Brochner- Mortensen, 1973)	89 clearances in 9 subjects	Healthy, before and after hyperglycemia	130 to 150	Urinary clearance and constant infused rate	Ratio	0.9±0.01
(Hagstam et al., 1974)	29	CKD	± 30 to 160	Urinary clearance and constant infused rate	Correlation Regression Ratio	0.97 =0.855x+7.555 0.96±0.07
(Hagstam et al., 1974)	31	CKD	± 30 to 160	Inulin: urinary clearance and constant infused rate 51Cr-EDTA: plasmatic clearance:	Correlation Regression Ratio	0.97 =0.961x+2.908 1±0.11
				samples at 180, 200, 220 et 240 min + BM correction		0.00
(Winterborn et al., 1977)	16	Children and 4 healthy adults	±5 to 120	Inulin: urinary clearance and constant infused rate ⁵¹ Cr-EDTA: urinary clearance:	Correlation Regression	0.99 =0.96x+3.5
(Jagenburg et al., 1978)	17	Severe CKD	2.6 to 11	Urinary clearance	Correlation Regression	0.97 =1.05x-0.3
(Rehling et al., 1986)	19	Nephrectomy	11 to 76	Inulin: urinary clearance and constant infused rate 51Cr-EDTA: plasmatic clearance: 5 samples between 3 and 5 h+BM correction	Correlation Regression SD around the mean difference	0.96 =0.86x+2.4 4.3 mL/min
(Froissart et al., 2005b)	111	NA	NA	Urinary clearance and constant infused rate	BA	2.7±3.5
(Froissart et al., 2005a)	22	NA	NA	Urinary clearance and constant infused rate	BA	4±4.9
(Medeiros et al., 2009)	44	Renal grafted	±15 to 80	Inulin: urinary clearance and constant infused rate ⁵¹ Cr-EDTA: plasmatic clearance: samples at 2, 4, 6, 8 h + BM correction	t-test Correlation BA Exactitude 30%	NS 0.94 2.5±6.1 90.9%

Table 2. Studies comparing ⁵¹Cr-EDTA with inulin. NA: not available, CKD: chronic kidney disease subjects, BA: Bland and Altman analysis, BAr: Bland and Altman analysis recalculated by us, BM: Brochner-Mortensen.

5.3 Strengths and limitations

⁵¹Cr-EDTA clearance was the first published alternative to inulin. Among the strengths of this marker, we have to underline the good performance of GFR measurement comparing to inulin (or to other markers). Physiological profile can also be considered as satisfying. This marker is yet easy to measure (especially according to its long half-life) and the precision of the measurement appears excellent. The costs, compared to other GFR markers, are acceptable. One important limitation is linked to the fact that ⁵¹Cr-EDTA GFR must be done in a Nuclear Medicine department. The most important limitation of this marker is the non-use in USA, where ⁵¹Cr-EDTA is not recognized by the FDA.

6. ⁹⁹Tc-DTPA (Diethylenetriaminepenta-acetic acid)

6.1 Physiological and analytical data

Like ⁵¹Cr-EDTA, ⁹⁹Tc-DTPA is an isotopic marker with a low molecular weight (393 Da)(Durand et al., 2006). DTPA may be labeled with another isotopic marker (^{113m}Indium (Johansson & Falch, 1978; Reba et al., 1968; Piepsz et al., 1974), ¹⁶⁹Ytterbium (Perrone et al., 1990; Russell et al., 1985)) but technetium 99 is the most used up to now. The ⁹⁹Tc-DTPA is also used in Nuclear Imagery (isotopic nephrogram) for instance to measure separately the function or the right and left kidney (Biggi et al., 1995; Hilson et al., 1976; Kainer et al., 1979). However, we will only discuss GFR measurement based on plasma and/or urinary methods with ⁹⁹Tc-DTPA. GFR can also be estimated with external counting using gamma camera (namely the "Gates" method) (Gates, 1984; Russell, 1987) but this method is not precise enough to be considered as a reference method for measuring GFR. For some authors, the GFR estimation given by the Gates method is even less performing than the creatinine clearance (Owen et al., 1982; Goates et al., 1990; van de Wiele C. et al., 1999; Ma et al., 2007; Mulligan et al., 1990; Galli et al., 1994; Ginjaume et al., 1985; Rodby et al., 1992; Tepe et al., 1987; Aydin et al., 2008; De Santo et al., 1999; Fawdry et al., 1985; Durand et al., 2006).

Doses of injected 99Tc-DTPA are totally safe (10 MBq)(Kempi & Persson, 1975; Durand et al., 2006). If the GFR measurement is coupled with nephrogram, the radioactive dose is however 40 to 200x higher than a simple GFR measurement with 51Cr-EDTA (Kempi & Persson, 1975; Griffiths et al., 1988). The half-life of 99Tc-DTPA is short (6.05 h) which imposes that the GFR measurement is realized quickly after the samplings, which is a practical inconvenient compared to 51Cr-EDTA (Owen et al., 1982). The 99Tc-DTPA measurement is as precise as other isotopic methods. The most relevant critic regarding 99Tc-DTPA is its potential binding to protein. This aspect has been debated in the literature. Some authors described a binding to plasma proteins from 2 to 13%, which implies an underestimation of GFR, especially when GFR is measured by plasmatic clearance (Kempi & Persson, 1975; Agha & Persson, 1977; Klopper et al., 1972; Biggi et al., 1995; Houlihan et al., 1999; Rehling et al., 2001). These high percentages could however been explained by the lack of purity of the first available preparations of 99Tc-DTPA (Rootwelt et al., 1980; Rehling et al., 2001; Fleming et al., 2004; Carlsen et al., 1980; Russell et al., 1983; Kempi & Persson, 1975). This hypothesis has been well illustrated in 1980 by Carlsen who studied and compared 51Cr-EDTA clearances with 4 different commercial preparations of 99Tc-DTPA. This author showed different results according to the preparation used (Carlsen et al., 1980). The binding to protein may also be studied by different methodologies (ultrafiltration, electrophoresis, precipitation, in vitro or in vivo, in humans or in animals etc)(Rehling et al.,

2001; Russell et al., 1983; Jeghers et al., 1990). For example, Rehling found a binding to protein of 10-13% but this author was also the only one who found a significant and comparable binding to protein for ⁵¹Cr-EDTA and iothalamate (Rehling et al., 2001). The subject is finally still debated (Jeghers et al., 1990). Another potential critic about ⁹⁹Tc-DTPA is the very poor available data on its physiological handling. A study in a dog model argued for the absence of tubular secretion and reabsorption (Klopper et al., 1972).

6.2 Clinical data

There are hopefully much more clinical studies comparing 99Tc-DTPA with other markers. After the preliminary study published by Hauser (Hauser et al., 1970), the performances of 99Tc-DTPA clearance was studied from the seventies. Klopper may be considered as one of the pioneers with this markers (Klopper et al., 1972). The first studies were however comparing 99Tc-DTPA with iothalamate and the samples were limited (Table 6)(Klopper et al., 1972; Rootwelt et al., 1980). The first study comparing 99Tc-DTPA with inulin was published in 1984 (Rehling et al., 1984). In table 3, we resumed the results of studies comparing 99Tc-DTPA with the gold standard method in adults. Two studies have compared with good statistical methods the urinary clearance of 99Tc-DTPA and inulin. In the study published by Lewis in 1989, the bias was excellent bias (near to 0) but the precision was not satisfying (± 18 mL/min)(Lewis et al., 1989). One year later, Perrone showed excellent concordance between urinary clearances of 99Tc-DTPA and inulin in 13 chronic kidney disease (CKD) patients. However, the results were less impressive in the 4 healthy subject where 99Tc-DTPA clearances overestimate (+12 mL/min) inulin clearances. Definitive conclusion about the performance of 99Tc-DTPA plasmatic clearance is difficult to draw and we clearly need additional studies on this topic.

6.3 Strengths and limitations

⁹⁹Tc-DTPA presents the advantages and inconvenient of other isotopic methods (see ⁵¹Cr-EDTA paragraph). The dosage of the marker is relatively cheap and precise. His short half-time makes it a few less practicable than ⁵¹Cr-EDTA. Among the most important advantages of ⁹⁹Tc-DTPA, we underline the fact that it is the only marker that can be coupled with nephrogram to give separated function between the two kidneys (Durand et al., 2006). Physiological data to confirm its role as a reference marker are however clearly lacking. We also think that global performance of ⁹⁹Tc-DTPA compared to inulin is probably a few less than the ⁵¹Cr-EDTA, especially with plasma clearances (at least in part because ⁹⁹Tc-DTPA is binding to proteins).

7. lothalamate

7.1 Physiological and analytical data

Iothalamate is an ionic contrast product which was particularly used for urography. Iothalamate is derived from the tri-iodobenzoic acid. Its molecular weight is 637 Da (Schwartz et al., 2006) and it is freely distributed into the extracellular volume (Visser et al., 2008). From a historical point of view, iothalamate was not the first contrast agent used to measure GFR. Other derivates from tri-iodobenzoic acid had been tested at the end of the fifties. Diatrizoate (Hypaque) was proposed by some authors as a potential GFR marker because it is fully excreted by the kidneys (Meschan et al., 1963; Burbank et al., 1963; Stokes et al., 1962; Mcchesney & Hoppe, 1957). However, other authors suggested that

References	Sample	Population	GFR range (mL/min/	GFR methods	Statistics	Results
(Rehling et al., 1984)	20	Nephrectomy	1.73 m²) 11 to 76	Inulin: urinary and plasma clearance with bolus	Wilcoxon Ratio urinary	urine : p<0.05 plasma : p<0.05
				99Tc-DTPA: Urinary and plasma clearance: samples at 5, 10, 20, 40, 60, 90, 120, 150, 180, 210, 240, 270, 300 min	plasma ⁹⁹ Tc- DTPA and urinary clearance of inulin Correlation Regression	0.97 =0.93x+6.8
(Shemesh et al., 1985)	45	NP	±10 to 140	Inulin: urinary clearance and constant infused rate ⁹⁹ Tc-DTPA urinary clearance	Correlation Ratio	0,969 1,02±0,14
(Notghi et al., 1986)	37	Healthy and CKD	7 to 182	Inulin: urinary clearance and constant infused rate 99Tc-DTPA plasma clearance: samples at: 60 and 150 min	Correlation Regression	0.77 =0.94x+33.7
(Petri et al., 1988)	NA	Lupus	23 to 123	Inulin: urinary clearance and constant infused rate 99Tc-DTPA: urinary clearance with bolus	Correlation Regression r ²	0.96 =x+4.4 0.93
(Lewis et al., 1989)	29	10 heart grafted 11 renal grafted 10 donors	10 to 117	Inulin: urinary and plasma clearance with bolus 99Tc-DTPA: urinary clearance with bolus	Correlation Regression BAr	0.85 =0.84x+8.4 0±18

(Perrone et	13	CKD	±5 to 130	Inulin:	Wilcoxon or	P<0.001
al., 1990)				urinary	t-test	
				clearance and	Correlation	from 0.93 to
				constant		0.98
				infused rate	BA	Day 1
				⁹⁹ Tc-DTPA		+0.5±3
				urinary		Day 2
				clearance with		-2±3
				bolus		
	15 7			1)) %		
	4	Healthy)) (Inulin (day 1
						and 2):
		2 successive				108±14
		days				96±8
						99Tc-DTPA (day
						1 and 2)
						122±24
						108±17
(Wharton, III	18	Intensive care	2 to 69	Inulin:	Correlation	0.85
et al., 1992)		and CKD		urinary	Regression	=1.12x
				clearance and		
				constant		
				infused rate		
				⁹⁹ Tc-DTPA		
				urinary		
				clearance with		
				bolus		
(Gunasekera	15	NA	±25 to 160	Bolus and	Correlation	=0.98x-0.4
et al., 1996)				plasma	Regression	0.98
				clearance for		
				inulin and		
				⁹⁹ Tc-DTPA: 6		
				samples		
				within the		
				first hours, 3		
				or 4 samples		
				between 2 and		
				4 h		

Table 3. Studies comparing ⁹⁹Tc-DTPA with inulin. NA: not available, CKD: chronic kidney disease subjects, BA: Bland and Altman analysis, BAr: Bland and Altman analysis recalculated by us, BM: Brochner-Mortensen.

diatrizoate (as other derivates from tri-iodobenzoic acid) was secreted by renal tubules (Woodruff & Malvin, 1960; Harrow, 1956; Winter & Taplin, 1958). In 1961, Denneberg is the first to compare diatrizoate labeled with l¹³¹ and inulin in human (Denneberg et al., 1961). This author described a higher renal excretion and then confirmed that diatrizoate is secreted by renal tubules (Denneberg et al., 1961). Diatrizoate was still studied by some authors in the next years but the interest has definitively moved from diatrizoate to iothalamate (Burbank et al., 1963; Morris et al., 1965; Dalmeida & Suki, 1988; Owman & Olin, 1978; Donaldson, 1968).

As we will describe in the next paragraph, interest in iothalamate as a GFR marker has grown from the mid-sixties with the studies proposed by Sigman (Sigman et al., 1965a;

Sigman et al., 1965b). For this author, the binding of iothalamate to protein is less than 3% (Sigman et al., 1965b). Such result was confirmed by most of the authors thereafter (Anderson et al., 1968; Gagnon et al., 1971; Blaufox & Cohen, 1970; Prueksaritanont et Back et al., 1988b), except for Maher and Rehling (see 99Tc-DTPA chapter)(Rehling et al., 2001; Maher & Tauxe, 1969). Rapidly, Sigman has proposed to move from labeling with I131 to labeling with I125. I125 is actually more stable (Elwood & Sigman, 1967; Maher et al., 1971). I125-Iothalamate is thus an isotopic method which is precise and safe. The half-life of 125I is 60 days (Perrone et al., 1990). Physiological data on iothalamate have been published after the first clinical studies by Sigman. Iothalamate was then studied in aglomerular fishes and only 3% of injected iothalamate was found in urine. The absence of tubular secretion and reabsorption was confirmed in a dog model (Griep & Nelp, 1969). However, these reassuring results were not confirmed by Odlind in 1985. This author actually observed in rats a tubular secretion of iothalamate (comparing with 51Cr-EDTA and using inhibitors of tubular secretion). In the same view, Odlind described, in 6 healthy subjects, that iothalamate clearance overestimates inulin clearance and that this overestimation is reversible after inhibition of tubular secretion by probenecid (Odlind et al., 1985). In anephric patients, Cangiano described an extra-renal excretion of iothalamate that reached 4 to 8 mL/min. This extra-renal excretion fall to 0 after thyroid saturation by iodine (Cangiano et al., 1971). A potential limited extra-renal clearance of iothalamate was thus suggested in the thyroid. Evans described a clearance of iothalamate of 3.1±1.8 mL/min in 7 dialysis patients (among these, 5 were anuric). In animal models, a limited biliary excretion is suggested by some authors (Owman & Olin, 1978; Prueksaritanont et al., 1986). Comparing the total (i.e. plasma) and the renal clearance of iothalamate in healthy subjects, Back calculated the extra-renal clearance at 6 mL/min (Back et al., 1988b). In the same experience, Dowling calculated extra-renal clearance at 10 ml/ml, which was constant for all the GFR levels (sample of 26 patients)(Dowling et al., 1999). In this last study, the plasma clearance was measured until 180 min, which may be considered as too short (Dowling et al., 1999). Visser has also calculated the urinary excretion of iothalamate on 24 h and estimated the extra-renal excretion at 14±12% (Visser et al., 2008). Such values of extra-renal clearances are thus not so negligible, especially when it is considered in patients with severe CKD. Actually, the relative importance of this extra-renal clearance will be higher when the GFR is yet low (Visser et al., 2008).

Iothalamate is a safe product but, of course, it will be not used in subjects presenting a known "true" allergy to contrast products (Heron et al., 1984). Regarding the isotopic method, the radioactive dose got by the patient is also very low (lower than the dose got for thorax radiography)(Hall & Rolin, 1995; Bajaj et al., 1996).

Because its relatively low molecular weight, iothalamate is a good marker (just like ⁵¹Cr-EDTA) to be used in simplified protocols. Cohen was the first to use the bolus method instead of the constant rate infusing method in 1969 (Cohen et al., 1969). Several authors have showed that iothalamate could be used in plasma clearance (LaFrance et al., 1988; Welling et al., 1976; Back et al., 1988b; Gaspari et al., 1992) even if results are not fully comparable to urinary clearances (Agarwal et al., 2009). It must also be underlined that iothalamate is the only one marker which is frequently used with subcutaneous injection (Israelit et al., 1973). It had actually been shown that plasma iothalamate concentrations remain constant 60 to 90 min after a subcutaneous injection (so, equivalent to the constant

infusion rate method but much easier) (Israelit et al., 1973; Adefuin et al., 1976; Tessitore et al., 1979; Sharma et al., 1997).

Iothalamate can yet be measured by "cold" non-isotopic methods. The first "cold" dosage of iothalamate was proposed in 1975 by Guesry (Guesry et al., 1975). This author used fluorescent excitation analysis or X ray fluorescence (XRF), which will be also used for iohexol measurement (see below). In this technique, iodine atoms are ionized by americanum. When the iodine atom comes back to neutral status, it will emit X ray that will be then quantified (Guesry et al., 1975). Guesry found an excellent correlation between isotopic and XRF iothalamate measurement. Iothalamate concentration can also be determined by electrophoresis but, to the best of our knowledge, this technique is only used in the Mayo Clinic (Wilson et al., 1997). The most used methods to measure iothalamate are HPLC methods (Boschi & Marchesini, 1981). The HPLC method seems specific, sensible and reproducible (CV intra-day lower than 2% and CV inter-day lower than 6%) (Boschi & Marchesini, 1981; Prueksaritanont et al., 1984; Weber et al., 1985; Reidenberg et al., 1988; Back et al., 1988b; Gaspari et al., 1991; Dowling et al., 1998; Agarwal, 1998; Kos et al., 2000; Agarwal et al., 2003; Farthing et al., 2005; Bi et al., 2007). A new technique based on mass spectrometry has recently been proposed to measure iothalamate (Seegmiller et al., 2010). These authors have compared 51 GFR results given by this new technique and by electrophoresis. The results are excellent in term of correlation and bias (0.8%). The SD around the bias, namely the precision, is however less negligible at 13.7%. That means that 95% of the results measures in the same patient may vary from ± 28% according the way iothalamate has been measured. Iothalamate measurement remain very stable (for two months at room temperature and at -4 and -20°C and for 1 year at -80°C) (Weber et al., 1985; Seegmiller et al., 2010).

7.2 Clinical data

Iothalamate (Conray°) was used as GFR marker for the first time by Sigman from the New York University in 1965 (Sigman et al., 1965a; Sigman et al., 1965b). In these articles, Sigman used iothalamate labeled with 131I and compared its clearance with inulin clearance in 10 patients in the first publication (Sigman et al., 1965a) and in 16 in the second one (Sigman et al., 1965b). On this limited sample, Sigman described a ratio iothalamate/inulin near to 1, even though the ranges of this ratio are from 0.74 à 1 in the first study (Sigman et al., 1965a) and from 0.937 à 1.138 in the second one (Sigman et al., 1965b). These first interesting results were then confirmed by the same authors with 125I-iothalamate (Elwood & Sigman, 1967). Other authors published thereafter their own data comparing performance of inulin and iothalamate clearances. We resumed the results obtained in adults in Table 4. It is probably right to write that iothalamate has been the most studied GFR marker and the marker for which several comparisons to inulin exist. Other authors have confirmed the good performance of iothalamate urinary clearances, especially in CKD patients (Maher et al., 1971; Perrone et al., 1990; Skov, 1970). In healthy subjects, the results are however more questionable and iothalamate seems to overestimate inulin (+20 mL/min)(Perrone et al., 1990) although precision is not optimal ±11 mL/min, as illustrated in the study by Botev (Botev et al., 2011). Data regarding the performance of the iothalamate plasma clearance are less numerous but is seems that bias is acceptable. However, precision is not optimal, especially in higher GFR levels. Additional studies could be of interest for the plasmatic method (Agarwal, 2003; Mirouze et al., 1972).

References	Sample	Population	GFR range (mL/min/1.73	GFR methods	Statistics	Results
			m ²)			
(Sigman et	10	NA	70 to 108	Inulin:	Ratio	1.06
al., 1965a)				urinary	It/inulin	(0.74 to 1.23)
				clearance and	BAr	6±13
				constant		
				infused rate		
	\neg \vdash \vdash			¹³¹ iothalamate		
	7 17 6			: urinary	$\bigcirc \bigcirc \bigcirc \bigcirc$	
			$\neg \Box \Box \Box \Box$	clearance and))(—	
				constant		
				infused rate		
(Sigman et	24	NA	2 to 167	Inulin:	t-test	NS
al., 1965b)	clearances			urinary	Ratio	1.005 (from 0.937
	in 16			clearance and	It/inulin	to 1.138)
	subjects			constant	BAr	0.7 ± 4
				infused rate		
				¹³¹ iothalamate		
				: urinary		
				clearance and		
				constant		
(5)				infused rate		
(Elwood &	26	NA	27 to 136	Inulin:	Ratio	1 (from 0.93 to
0 '	clearances			urinary	It/inulin	1.09)
1967)	in 21			clearance and	BAr	1±3
	subjects			constant		
				infused rate		
				¹²⁵ iothalamate		
				: urinary		
				clearance and		
				constant		
/N /-1	10	Harlibara ACVD	NTA	infused rate	D.C.	1.01+0.10
(Malamos et	19	Healthy and CKD	NA	Inulin:	Ratio	1.01±0.19
al., 1967)				urinary	It/inulin	0.070
				clearance and		0.979
				constant	(urinary)	I+-1 (10):1: 0 (5
	\nearrow \vdash			infused rate ¹²⁵ iothalamate	Regression	It=1.09inulin-0.65
				: urinary		
			$\neg \sqcap \sqcap \sqcap $	clearance and	$\bigcup \setminus \setminus \subset$	71111
				constant		
				infused rate		
(Anderson et	18	11 CKD and 8	3 to 139	Inulin:	Regression	=0.9x+6.7
al., 1968)	10	healthy	J 10 139	urinary	BAr	-0.7±13
(1., 1700)		riculary		clearance and	<i>D1</i> 11	0.7 ±10
				constant		
				infused rate		
				¹²⁵ iothalamate		
				: urinary		
				clearance and		
				constant		
				infused rate		

(Maher &	15	hypertensive	±55 to 120	Inulin:	Ratio	0.92
Tauxe, 1969)	15	Try per terisive	±55 to 120	urinary	It/inulin	(0.81 to 1.04)
,				clearance and	Regression	Inulin=1.08It
				constant		
				infused rate		
				¹²⁵ iothalamate		
				: urinary		
				clearance and constant		
				infused rate		
(Skov, 1970)	43	CKD CKD		Inulin:		Group 1
			\neg $ $ $ $ $ $ $ $	urinary	Ratio	0.98 ± 0.06
	65	GFR<5 ml/		clearance and	It/inulin	
	clearances			constant	Correlation	0.999
	in 22			infused rate	Regression	=0.972+0.01
	subjects	GFR between 5 et		¹²⁵ iothalamate	BAr	0±0
	38	15 mL/min		: bolus and		Group 2
		GFR between 15 et		urinary clearance		1 0±1
	in 13	25 mL/min		Clearance		0±1
	subjects					Group 3
						0.92±0.071
						0.968
	24					=1.083+3.46
	clearances					-2±1
	in 8					
(Company of	subjects 78	NA	±10 to 180	Inulin:	Ratio	1.01
(Gagnon et al., 1971)	78 clearances	NA	±10 to 180	urinary	Ratio It/inulin	1.01
ai., 1971)	in 24			clearance and	n, mam	
	subjects			constant		
	Surjects			infused rate		
				¹²⁵ iothalamate		
				: urinary		
				clearance and		
				constant		
(Congione et	40	NA	±30 to 150	infused rate	Ratio	1.07
(Cangiano et al., 1971)	49 clearances		±30 to 150	Inulin: urinary	Ratio It/inulin	1.0/
ai., 19/1)	in 18			clearance and	Correlation	0.94
	subjects			constant	Regression	=1.06+1.17
	75-			infused rate		
				¹²⁵ iothalamate))((
			71111	: urinary		71111
				clearance and		
				constant		
(Maher et al.,	198	NA	±5 to 150	infused rate Inulin:	Bias	-2.09
1971)	190	INA	±3 tO 130	urinary	Dias	-2.09
17/1)				clearance and	Regression	Inulin=1.022It+0.5
				constant	-1062 0001011	37
				infused rate		
				¹²⁵ iothalamate		
				: urinary		
				clearance and		
				constant		
				infused rate		

(Mirouze et	36	hypertensive	±5 to 120	Inulin:	Ratio	1.44±0.13
al., 1972)	clearances	ny pertensive		urinary	It/inulin	11110110
, ,	in 23			clearance and	•	0.96
	subjects			constant	Regression	=1.18+8.43
	,			infused rate	Ü	
				¹²⁵ iothalamate		
				: urinary		
				clearance and		
				constant		
	7) 7 [infused rate	$\bigcirc \bigcirc \bigcirc \bigcirc$	
(Mirouze et	15	hypertensive	±80 to 140	Inulin:	Ratio	1.23±0.16
al., 1972)				urinary	It/inulin	
				clearance and	Correlation	0.77
				constant	Regression	=1.06+1.18
				infused rate		
				125iothalamate		
				:		
				plasma		
				clearance:		
				samples at 5,		
				10, 15, 20, 40,		
				60, 80, 100 et		
				120 min +		
				correction		
(Israelit et al.,	22	20 CKD	6 to 125	Inulin:	Ratio	1.05±0.04
1973)		2 healthy		urinary	It/inulin	
		-		clearance and	Correlation	0.97
				constant	Regression	=1.054-3.069
				infused rate		
				¹²⁵ iothalamate		
				: bolus SC and		
				urinary		
				clearance		
(Rosenbaum	7 healthy		96 to 147	Inulin:	Ratio	1.02±0.04
et al., 1979)	9 renal		35 to 87	urinary	It/inulin	1.43±0.08
	grafted		42 to 98	clearance and		1.23±0.04
	8 donors			constant	BAr	-1±13
	after	$(\triangle)(\triangle)$		infused rate	$\bigcirc \bigcirc \bigcirc \bigcirc$	-7±14
	donation			¹²⁵ iothalamate))((-4±13
				: bolus and		
				urinary		
				clearance		
(Ott, 1975)	84	CKD and donors	±10 to 150	Inulin:	Correlation	0.932
				urinary	Regression	=1.04+2.11
				clearance and		
				constant		
				infused rate		
				¹²⁵ iothalamate		
				: urinary		
				clearance and		
				constant		
				infused rate		

(Ott, 1975) (Tessitore et al., 1979)	30	15 creatinine<1 mg/dL 15 creatinine<20 mg/dL	±5 to 150	Inulin: urinary clearance and constant infused rate 125iothalamate bolus SC and urinary clearance Inulin: urinary clearance and constant	Ratio It/inulin Correlation	0.982 =1.02-0.61 1.07±0.05 0.96
		8, 1		infused rate 125iothalamate : bolus SC and urinary clearance		
(Notghi et al., 1986)	clearances in 40 subjects	Healthy and CKD	±10 to 180	Inulin: urinary clearance and constant infused rate 125 iothalamate bolus SC and urinary clearance	Correlation Regression	0.86 =0.8x+19.5
(Petri et al., 1988)	NA	Lupus	23 to 123	Inulin: urinary clearance and constant infused rate Iothalamate (XRF): bolus and urinary clearance	Correlation Regression r ²	0.99 =0.9x-2.1 0.99
(Perrone et al., 1990)	13	CKD	±5 to 130	Inulin: urinary clearance and constant infused rate	Wilcoxon or t-test Correlation	P<0.001 from 0.93 to 0.98
	4	Healthy Two successive days		125iothalamate : bolus SC and urinary clearance	Means	Inulin: 108±14 day 1 96±8 day 2 ¹²⁵ iothalamate 127±12 day 1 120±7 day 2
(al Uzri et al., 1992)	5	healthy	120 to 165	Inulin: urinary clearance and constant infused rate Iothalamate (HPLC): bolus and urinary clearance	ratio	1.00±0.06

(Isaka et al., 1992)	23	CKD	10 to 130	Inulin: urinary clearance and constant infused rate Iothalamate (HPLC): bolus and urinary clearance	Correlation Slope with 0 intercept	0.98 1.05±0.01
(Agarwal, 2003)	12 clearances in 3 subjects	CKD	± 20 to 110	Inulin: urinary clearance and constant infused rate Iothalamate (HPLC): plasma clearance on long time with insulin pomp	Bias (Inulin- It) CV	0.8 19.9%
(Botev et al., 2011) Data from 5 studies (Anderson et al., 1968; Elwood & Sigman, 1967; Perrone et al., 1990; Rosenbaum et al., 1979; Skov, 1970)		See above	±5 to 140	See above	Correlation Regression BA (It-Inulin)	0.97 =1.04+2.334 +4.6±11

Table 4. Studies comparing iothalamate with inulin. NA: not available, CKD: chronic kidney disease subjects, BA: Bland and Altman analysis, BAr: Bland and Altman analysis recalculated by us, BM: Brochner-Mortensen, HPLC: high pressure liquid chromatography, It: iothalamate, SC: subcutaneous, XRF: X ray fluorescence.

7.3 Strengths and limitations

Iothalamate can be measured either by HPLC or XRF methods or by isotopic methods. This is the only one marker where this choice is possible. However, there is no evidence that all the techniques of measurement are fully equivalent. Iothalamate is certainly the marker that has been the most deeply studied from a physiological point of view (with inulin). Unhopefully, there are strong reasons to believe that iothalamate is secreted by renal tubules. Moreover, extra-renal clearance of iothalamate is not so negligible. These limitations are confirmed by most of the clinical studies showing that iothalamate slightly overestimates inulin clearance, especially in the high levels of GFR. A clinical limitation concerns the patients who are allergic to contrast product. This marker remains however

important because it is the most used marker in USA. For example, iothalamate has been used in trials having built the new creatinine-based equations (Levey et al., 1999).

8. lohexol

8.1 Physiological and analytical data

Iohexol is a non-ionic contrast product, mainly used for myelography. Its molecular weight is 821 Da (Olsson et al., 1983; Schwartz et al., 2006). Iohexol is chronologically the last marker proposed for measuring GFR. Actually, the first human was receiving iohexol in 1980 (Aakhus et al., 1980). In this study, it was shown that the substance was safe and fully excreted by the kidneys (this assertion will be criticized thereafter, see below). However, these authors also describe (but data are not available) a higher urinary clearance of iohexol than 51Cr-EDTA (Aakhus et al., 1980). The details of these comparison studies were published three years after (see clinical data)(Olsson et al., 1983). In the same study, the authors confirm that iohexol is distributed through the extracellular volume, which will be confirmed by other authors (including in CKD patients and in obese subjects) (Friedman et al., 2010; Nossen et al., 1995; Edelson et al., 1984; Back et al., 1988b; Olsson et al., 1983). Iohexol has not effect per se on GFR (Olofsson et al., 1996). Binding to protein seems very limited for iohexol. The first study described a binding to protein of only 1.5% (Mutzel et al., 1980). This will be thereafter confirmed (Back et al., 1988b; Krutzen et al., 1984). Physical properties of iohexol make it a good candidate to be used in simplified protocols like plasma clearance (Thomsen & Hvid-Jacobsen, 1991; Gaspari et al., 1995; Edelson et al., 1984). Contrary to the prior studies (Aakhus et al., 1980), several authors have shown that extrarenal clearance of iohexol is limited but not null (Arvidsson & Hedman, 1990; Krutzen et al., 1984). Back calculated at 6.2 mL/min the difference between total and urinary clearance of iohexol in healthy subjects (Back et al., 1988b). Frennby observed an extra-renal clearance lower than 2 mL/min in 6 anuric dialysis patients (Frennby et al., 1994; Frennby et al., 1995). These last very low results were also found by Nossen in 16 patients with severe CKD. Their mean measured GFR was 14 mL/min and the extra-renal clearance was estimated at 10% (Nossen et al., 1995). In 16 healthy subjects, Edelson estimated the extrarenal clearance of iohexol at 5% (Edelson et al., 1984). Contrary to iothalamate, there are very few physiological studies on the renal tubular handling of iohexol.

As for iothalamate, iohexol can be measured by several different techniques. Among these, HPLC and XRF are the most used ones. HPLC was historically the first method used (Aakhus et al., 1980) and described (Krutzen et al., 1984). As we have shown, iohexol measurements by HPLC are sensitive, specific and reproducible (Back et al., 1988c; Farthing et al., 2005; Cavalier et al., 2008). The high performance of such dosage notably enables the use of iohexol low doses and the measurement on finger-prick samples (Krutzen et al., 1990; Niculescu-Duvaz et al., 2006; Mafham et al., 2007; Cavalier et al., 2008; Aurell, 1994). Iohexol measurement is also pretty stable at room temperature and at -20°C(Krutzen et al., 1984; O'Reilly et al., 1988). Measurement of iohexol by XRF method is less validated and probably less performing, especially in low plasma concentrations (O'Reilly et al., 1986; Back & Nilsson-Ehle, 1993; Effersoe et al., 1990; Brandstrom et al., 1998; Aurell, 1994). We will not discuss into details the other methods for measuring iohexol: capillary electrophoresis (Shihabi & Constantinescu, 1992) and mass spectrometry (Lee et al., 2006; Annesley & Clayton, 2009; Denis et al., 2008; Stolz et al., 2010). The safety of iohexol is now confirmed (Heron et al., 1984; Aurell, 1994), notably by the largest series of iohexol

measurements in Sweden (1500 GFR measurements/y)(Nilsson-Ehle & Grubb, 1994; Nilsson-Ehle, 2002). This safety profile is, at least in part, explained by the low dose of iohexol injected, and by the exclusion of patients with contrast products allergy.

8.2 Clinical data

The results of the first clinical study on iohexol as a reference GFR marker will be published in 1983 (Olsson et al., 1983). Actually, GFR was measured in 10 healthy subjects with urinary clearances of iohexol and 51Cr-EDTA. In this study, the iohexol clearance was significantly higher than the 51Cr-EDTA clearance (110 versus 96 mL/min). In this first study, large dose of iohexol was injected to the patient (from 375 to 500 mg I/kg)(Olsson et al., 1983). Thereafter, the doses of iohexol used will be drastically reduced but it has been well described that the physiologic handling of iohexol was identical if different dosages are used (Back et al., 1988a). In table 5, we resumed the study results having compared the performance of iohexol to inulin in adult subjects. To the best of our knowledge, only two studies have compared urinary clearances of iohexol and inulin. The results seem excellent but Bland and Altman analysis have not been realized (Brown & O'Reilly, 1991; Perrone et al., 1990). Contrary to other markers, iohexol plasmatic clearances have been the most studied. The relatively worst results obtained by Erley are explained by the patients included (Erley et al., 2001). Actually, the patients hospitalized in intensive care are prone to develop edema and, in this situation, plasmatic clearances are not accurate, whatever the marker (Skluzacek et al., 2003). The study published by Gaspari demonstrated a good performance of iohexol plasma clearance compared to inulin but the number of samples was high and these samples were drawn lately (after 10h)(Gaspari et al., 1995).

References	Sample	Population	GFR range	GFR methods	Statistics	Results
			(mL/min/1.73			
			m²)			
(Lewis et al.,	29	10 heart	9.6 to 116.8	Inulin: urinary	Correlation	0.86
1989)		grafted		clearance and	Regression	=0.85x+8.79
		11 renal grafted		constant	Ratio	1.09±0.06
		10 donors		infused rate		
				Iohexol (XRF)		
				Plasma		
				clearance:		
				bolus and		
				samples after 3)) (<u> </u>
			711 11\	and 4		7
(Brown &	30	NA	±10 to 125	Inulin: urinary	Correlation	Urinary
O'Reilly,				clearance and	Regression	0.986
1991)				constant	Ratio	=0.998-2.309
				infused rate		Plasma
				Iohexol (XRF)		0.983
				urinary		=0.947+4.92
				clearance and		=1.102±0.286
				plasma		
				clearance:		
				samples at 3		
				and 4 h +BM		
				correction		

(Gaspari et al., 1995)	41	CKD	6 to 160	Inulin: urinary clearance and constant infused rate Iohexol (HPLC) Plasma clearance, samples at 5, 10, 20, 30, 45, 60, 90, 120, 180, 240, 300, 450, 600 min	Correlation Regression BA	0.97 =0.994x+2.339 1.02±7
(Erley et al., 2001)	31	intensive care	±10 to 130	Inulin: urinary clearance and constant infused rate Iohexol (XRF) Plasma clearance: samples at 150, 195, 240 + 360 min if estimated GFR under 30 mL/min	Correlation Regression BA	=0.971x+7.65 r ² =0.96 (Io-inulin) =8.67±7.21
(Sterner et al., 2008)	20	healthy	106 to 129	Inulin: urinary clearance and constant infused rate Iohexol (HPLC) Urinary clearance and constant infused rate	Wilcoxon	Not different

Table 5. Studies comparing iohexol with inulin. NA: not available, CKD: chronic kidney disease subjects, BA: Bland and Altman analysis, BAr: Bland and Altman analysis recalculated by us, BM: Brochner-Mortensen, HPLC: high pressure liquid chromatography, Io: iohexol, SC: subcutaneous, XRF: X ray fluorescence.

8.3 Strengths and limitations

Iohexol is probably the easiest way to measure GFR. It can be used in all patients (except in patient with true allergy to contrast product). Its measurement by HPLC is probably one of the most precise compared to other cold method (inulin and iothalamate). Iohexol is the less expensive marker and the cost of HPLC is also low. More important, it must be underlined that an external quality control does exist for iohexol measurement (Equalis, Sweden). From unpublished data, it can be concluded that the inter-laboratory CV for iohexol measurement is very low (less than 5%). Such results don't exist for iothalamate and inulin, and, at least for inulin, we think that such good inter-laboratory results would not be reached (personal

data). The limitations of iohexol are the lack of strong physiological data (notably regarding the tubular handling of the marker) and the relatively few studies having compared iohexol with inulin. More studies have actually compared iohexol with other GFR markers.

9. Studies comparing reference methods

In Table 6, we resumed the results of studies comparing reference markers (other than inulin). We selected studies in adults. We focused on studies having used the best statistical methods to analyze the results, i.e. the Bland and Altman analysis. It is difficult to interpret results from studies having compared different markers but also different methods (for example, plasmatic clearance of iothalamate with urinary clearance of ⁵¹Cr-EDTA) because it is impossible to affirm that potential differences are due to difference in markers or to difference in methods. Another limitation of several studies is the relatively small sample of subjects included. If we take into account these two limitations, we can stress on some interesting results showing good concordance (bias±SD) between plasma clearances of ⁵¹Cr-EDTA and ⁹⁹Tc-DTPA (1.91±6.1 mL/min), and between plasma clearances of ⁵¹Cr-EDTA and iohexol (-0.16±6.17 mL/min in (Brandstrom et al., 1998), 4±7.9 mL/min in (Bird et al., 2009), 2±9.2 (Lundqvist et al., 1997), and -0.6±3.6 mL/min in (Pucci et al., 2001)).

References	Sample	Population	GFR range (mL/min/1.73 m²)	GFR methods	Statistics	Results
(Odlind et al., 1985)	11	Nephrectomy and CKD	37 to 137	Cp of Cr and 125It: samples at 180, 210 and 240 min + BM correction	Wilcoxon Ratio It/Cr BAr It-Cr	It higher (p<0.001) 1.13 12±7.5
(Lewis et al., 1989)	29	10 heart grafted 11 renal grafted 10 donors	10 to 117	Cu of Dt and Io (XRF): samples at 3 and 4 h after bolus	Correlation Regression Ratio BAr Dt-Io	0.89 Io=0.89Dt+6.5 1.08±0.06 -0.7±14.8
(Goates et al., 1990)	16	NA C	21 to 156	Cu ¹²⁵ It: Cu after bolus IV and infusion Cp of Dt: samples at 60 and 180 min+ BM correction	Correlation BAr Io-Dt	0.99 3.2±6.1
(Effersoe et al., 1990)	15	urography	22 to 110	Cp of Io (XRF), Cr and Dt: samples at 0, 10, 20, 30, 120, 180, 240 and 300 min	Regression Correlation BA : Cr-Io Dt-Io Cr-Dt	Io=0.97Dt-11 0.96 Io=1.01Cr+8 0.95 -10.8±7.9 -9.4±6.9 -0.7±10.4

(Gaspari et al., 1992)	19	CKD	7 to 148	Cp of Cr and It (HPLC):	Correlation Regression BAr It-Cr	0.995 It=1.007Cr-0.303 -0.1±4.7
				bolus IV and samples at 5,	DAI II-CI	-U.11 1 ./
				10, 20, 30, 40, 50, 60, 90, 120,		
				180, 240, 300,		
				450 and 600		
(Lundariet et	31	Para or	±70 to 130	min Cn of Cr and	BA Cr-Io	Day 1 . +2 1+10 2
(Lundqvist et al., 1994)	31	tetraplegic	±70 to 130	Cp of Cr and Io (XRF):	DA CI-10	Day 1 : +2.1±10.2 Day 2 : +0.9±5.9
*()				samples à 180,		,
				210, 240 and		
				270 minutes+		
				BM correction		
(0.11)	=0	774	15.10	day 1 and 2		D. 0000G
(Galli et al., 1994)	50	NA	±15 to 160	Cp of Cr and	Regression BA Dt-Cr	Dt=0.982Cr+3.2 1.91±6.1
1994)				Dt: samples at 60 and 180	DA DI-CI	1.91±0.1
				min		
(Sambataro	17	Diabetic	7 to 105	Cu of Cr and	Regression	It=0.979Cr-3.04
et al., 1996)				It (HPLC)	BAr It-Cr	1.3±5
(Lundqvist et	77	Urography	±25 to 125	Cp of Cr and	Correlation	0.918
al., 1997)				Io (XRF):	Regression	Io=0.892Cr+6.28
				samples at	BA (Io-Cr)	2±9.2
				180, and 240 or 270		
				min+ BM		
				correction		
(Brandstrom	49	GFR>40	±40 to 125	Cp of Cr and		XRF
et al., 1998)				Io (HPLC and	Regression	Io=1.03Cr-1.79
				XRF): samples	Correlation BA Cr-Io	0.97
				at 150, 195	BA Cr-10	0.58±4.95 HPLC
				and 240 min + BM correction		Io=1.05Cr-4.43
	1)77/		7101	Divi correction		0.96
(D) : 1	22		10: 151			-0.16±6.17
(Pucci et al., 1998)	32	Diabetic	13 to 151	Cp of Cr and Io (HPLC):	Regression Correlation	0.995 Io=0.978Cr+2.45
1990)				samples at 5,	BA Cr-Io	-0.6±3.6
				10, 15, 30, 60,		
				90, 120, 150,		
				180, 210, 240,		
				270, 300 + 360		
				and 420 if		
				creatinine>2		
				mg/dL + 1440 min		
				if>5mg/dL		
				11/JIIIg/ UL		

Houlihan	21	Diabetic	50 to 145	Cp of Dt and	Regression	Io=0.9938Dt+4.91
(Houlihan et				Io (XRF):	Correlation	6
al., 1999)				samples at	BA Io-Dt	0.97
ŕ				120, 165 and		4.3±7.7
				210 for Dt		
				samples at		
				120, 150, 180,		
				210 and 240		
				min for Io		
				+ BM		
			$\neg \square \square \square$	correction		
(Pucci et al.,	41	Diabetic	29 to 150	Cp of Cr and		Type 1
2001)				Io (HPLC):	Regression	Io=0.978Cr+0.132
				samples at 5,	Correlation	0.999
				10, 15, 30, 60,		Type 2
				90, 120, 150,		0.987
				180, 210, 240,		Io=0.078Cr+2.352
				270, 300 + 360		
				and 420 if	BA Cr-Io	BA :-0.42±3.69
				creatinine>2		
				mg/dL+		
				1440 min		
				if>5mg/dL		
Bird (Bird et	56	CKD	±15 to 140	Cp of Cr and	BA Cr-Io	4±7.9
al., 2009)				Io (XRF):		
	19	healthy		samples at 20,		
				40, 60, 120,		
				180 and 240		
				min		

Table 6. studies comparing different reference methods (other than inulin), NA = not available, BA: Bland and Altman, BAr: Bland and Altman recalculated by us, BM: Brochner Mortensen,Cr: ⁵¹Cr-EDTA, Dt: ⁹⁹Tc-DTPA, Io: iohexol, It: iothalamate, Cp: plasma clearance, Cu: urinary clearance, IC: constant infusion rate, IB: bolus injection, IV: intravenous, SC: subcutaneous, AUC: area under the curve, NS: not significant, HPLC: high pressure liquid chromatography, XRF: X ray fluorescence.

10. Conclusions and perspectives

In this chapter, we reviewed all the reference methods available in 2011 to measure GFR. Among these methods, inulin clearance can certainly be considered as the gold-standard because it is historically the first method used and because this marker is certainly the best characterized from a physiological point of view. However, inulin is expensive and commercial sources are limited (Gaspari et al., 1997). Due to its high molecular weight, there are doubts to use inulin in simplified plasma clearance (urinary clearances with constant infusion rate remain necessary but are very cumbersome). Measurement of plasma inulin is neither easy nor standardized. For all these reasons, the use of inulin is and will always be relatively marginal. In 2011, it is maybe time to move from the perfect physiological marker (inulin) to markers, maybe less perfect in the renal physiologic handling, but less costly, easier to use everywhere in the world and with a standardized measurement. From our point of view, iohexol is probably the best marker with the best balance between

physiological characteristics and practical advantages. Additional studies comparing references markers seem necessary in 2011. It seems also important to underline that GFR measurement is also subject to its own imprecision and to biological variation (Kwong et al., 2010). Therefore, it is illusionary to expect differences between different GFR methods of less than 10% (±2SD around the bias) and accuracy 10% over 85-90%. We must also keep these results in mind when we analyze the studies testing the performance of the creatinine-based equations (Kwong et al., 2010).

11. References

- Aakhus, T., Sommerfelt, S.C., Stormorken, H. & Dahlstrom, K. (1980). Tolerance and excretion of iohexol after intravenous injection in healthy volunteers. Preliminary report, *Acta Radiol Suppl*, Vol.362, pp. 131-134
- Adefuin, P.Y., Gur, A., Siegel, N.J., Spencer, R.P. & Hayslett, J.P. (1976). Single subcutaneous injection of iothalamate sodium I 125 to measure glomerular filtration rate, *JAMA*, Vol.235, No.14, pp. 1467-1469
- Agarwal, R. (1998). Chromatographic estimation of iothalamate and p-aminohippuric acid to measure glomerular filtration rate and effective renal plasma flow in humans, *J Chromatogr B Biomed Sci Appl*, Vol.705, No.1, pp. 3-9
- Agarwal, R. (2003). Ambulatory GFR measurement with cold iothalamate in adults with chronic kidney disease, *Am J Kidney Dis*, Vol.41, No.4, pp. 752-759
- Agarwal, R., Bills, J.E., Yigazu, P.M., Abraham, T., Gizaw, A.B., Light, R.P., Bekele, D.M. & Tegegne, G.G. (2009). Assessment of iothalamate plasma clearance: duration of study affects quality of GFR, *Clin J Am Soc Nephrol*, Vol.4, No.1, pp. 77-85
- Agarwal, R., Vasavada, N. & Chase, S.D. (2003). Liquid chromatography for iothalamate in biological samples, *J Chromatogr B Analyt Technol Biomed Life Sci*, Vol.785, No.2, pp. 345-352
- Agha, N., Persson, R.B. (1977). Comparative labelling and biokinetic studies of 99mTc-EDTA(Sn) and 99mTc-DTPA(Sn), *Nuklearmedizin*, Vol.16, No.1, pp. 30-35
- al Uzri, A., Holliday, M.A., Gambertoglio, J.G., Schambelan, M., Kogan, B.A. & Don, B.R. (1992). An accurate practical method for estimating GFR in clinical studies using a constant subcutaneous infusion, *Kidney Int*, Vol.41, No.6, pp. 1701-1706
- Alving, A.S., Miller, B.F. & Rubin, J. (1939). A direct colorimetric method for the determination of inuline in blood and urine, *J Biol Chem*, Vol.127, No.3, pp. 609-616
- Anderson, C.F., Sawyer, T.K. & Cutler, R.E. (1968). Iothalamate sodium I 125 vs cyanocobalamin Co 57 as a measure of glomerular filtration rate in man, *JAMA*, Vol.204, No.8, pp. 653-656
- Annesley, T.M., Clayton, L.T. (2009). Ultraperformance liquid chromatography-tandem mass spectrometry assay for iohexol in human serum, *Clin Chem*, Vol.55, No.6, pp. 1196-1202
- Arvidsson, A., Hedman, A. (1990). Plasma and renal clearance of iohexol--a study on the reproducibility of a method for the glomerular filtration rate, *Scand J Clin Lab Invest*, Vol.50, No.7, pp. 757-761
- Aurell, M. (1994). Accurate and feasible measurements of GFR--is the iohexol clearance the answer?, *Nephrol Dial Transplant*, Vol.9, No.9, pp. 1222-1224
- Aydin, F., Gungor, F., Cengiz, A.K., Tuncer, M., Mahsereci, E., Ozdem, S., Cenkci, M. & Karayalcin, B. (2008). Comparison of glomerular filtration rate measurements with

- the two plasma sample and single plasma sample, gamma camera Gates, creatinine clearance, and prediction equation methods in potential kidney donors with normal renal function, *Nucl Med Commun*, Vol.29, No.2, pp. 157-165
- Baccard, N., Hoizey, G., Frances, C., Lamiable, D., Trenque, T. & Millart, H. (1999). Simultaneous determination of inulin and p-aminohippuric acid (PAH) in human plasma and urine by high-performance liquid chromatography, *Analyst*, Vol.124, No.6, pp. 833-836
- Back, S.E., Krutzen, E. & Nilsson-Ehle, P. (1988a). Contrast media and glomerular filtration: dose dependence of clearance for three agents, *J Pharm Sci*, Vol.77, No.9, pp. 765-767
- Back, S.E., Krutzen, E. & Nilsson-Ehle, P. (1988b). Contrast media as markers for glomerular filtration: a pharmacokinetic comparison of four agents, *Scand J Clin Lab Invest*, Vol.48, No.3, pp. 247-253
- Back, S.E., Masson, P. & Nilsson-Ehle, P. (1988c). A simple chemical method for the quantification of the contrast agent iohexol, applicable to glomerular filtration rate measurements, *Scand J Clin Lab Invest*, Vol.48, No.8, pp. 825-829
- Back, S.E., Nilsson-Ehle, P. (1993). Re: Iohexol clearance for the determination of glomerular filtration rate in clinical practice: evidence for a new gold standard, *J Urol*, Vol.149, No.2, pp. 378
- Bailey, R.R., Rogers, T.G. & Tait, J.J. (1970). Measurement of glomerular filtration rate using a single injection of 51Cr-Edetic acid, *Australas Ann Med*, Vol.19, No.3, pp. 255-258
- Bajaj, G., Alexander, S.R., Browne, R., Sakarcan, A. & Seikaly, M.G. (1996). 125Iodine-iothalamate clearance in children. A simple method to measure glomerular filtration, *Pediatr Nephrol*, Vol.10, No.1, pp. 25-28
- Bi, D., Leary, K.J., Weitz, J.A., Cherstniakova, S.A., Reil, M.A., Roy, M.J. & Cantilena, L.R. (2007). High performance liquid chromatographic measurement of iothalamate in human serum and urine for evaluation of glomerular filtration rate, *J Chromatogr B Analyt Technol Biomed Life Sci*, Vol.856, No.1-2, pp. 95-99
- Biggi, A., Viglietti, A., Farinelli, M.C., Bonada, C. & Camuzzini, G. (1995). Estimation of glomerular filtration rate using chromium-51 ethylene diamine tetra-acetic acid and technetium-99m diethylene triamine penta-acetic acid, *Eur J Nucl Med*, Vol.22, No.6, pp. 532-536
- Bird, N.J., Peters, C., Michell, A.R. & Peters, A.M. (2009). Comparison of GFR measurements assessed from single versus multiple samples, *Am J Kidney Dis*, Vol.54, No.2, pp. 278-288
- Bland, J.M., Altman, D.G. (1986). Statistical methods for assessing agreement between two methods of clinical measurement, *Lancet*, Vol.1, No.8476, pp. 307-310
- Blaufox, M.D., Cohen, A. (1970). Single-injection clearances of iothalamate-131-I in the rat, *Am J Physiol*, Vol.218, No.2, pp. 542-544
- Boschi, S., Marchesini, B. (1981). High-performance liquid chromatographic method for the simultaneous determination of iothalamate and o-iodohippurate, *J Chromatogr*, Vol.224, No.1, pp. 139-143
- Botev, R., Mallie, J.P., Wetzels, J.F., Couchoud, C. & Schuck, O. (2011). The Clinician and Estimation of Glomerular Filtration Rate by Creatinine-based Formulas: Current Limitations and Quo Vadis, *Clin J Am Soc Nephrol*, Vol.6, No.4, pp. 937-950

- Brandstrom, E., Grzegorczyk, A., Jacobsson, L., Friberg, P., Lindahl, A. & Aurell, M. (1998). GFR measurement with iohexol and 51Cr-EDTA. A comparison of the two favoured GFR markers in Europe, *Nephrol Dial Transplant*, Vol.13, No.5, pp. 1176-1182
- Brochner-Mortensen, J. (1973). The glomerular filtration rate during moderate hyperglycemia in normal man, *Acta Med Scand*, Vol.1-2, No.1, pp. 31-37
- Brochner-Mortensen, J. (1978). Routine methods and their reliability for assessment of glomerular filtration rate in adults, with special reference to total [51Cr]EDTA plasma clearance, *Dan Med Bull*, Vol.25, No.5, pp. 181-202
- Brochner-Mortensen, J., Giese, J. & Rossing, N. (1969). Renal inulin clearance versus total plasma clearance of 51Cr-EDTA, *Scand J Clin Lab Invest*, Vol.23, No.4, pp. 301-305
- Brochner-Mortensen, J., Rodbro, P. (1976). Comparison between total and renal plasma clearance of [51Cr] EDTA, *Scand J Clin Lab Invest*, Vol.36, No.3, pp. 247-249
- Brown, S.C., O'Reilly, P.H. (1991). Iohexol clearance for the determination of glomerular filtration rate in clinical practice: evidence for a new gold standard, *J Urol*, Vol.146, No.3, pp. 675-679
- Burbank, M.K., Tauxe, W.N., Maher, F.T. & Hunt, J.C. (1963). Utilisation des substances marquées dans les épreuves classiques de clearance rénale, *J Physiol (Paris)*, Vol.55, pp. 433-444
- Cangiano, J.L., Genuth, S.M., Renerts, L. & Berman, L.B. (1971). Simplified measurement of glomerular filtration rate, *Invest Urol*, Vol.9, No.1, pp. 34-38
- Carlsen, J.E., Moller, M.L., Lund, J.O. & Trap-Jensen, J. (1980). Comparison of four commercial Tc-99m(Sn)DTPA preparations used for the measurement of glomerular filtration rate: concise communication, *J Nucl Med*, Vol.21, No.2, pp. 126-129
- Cavalier, E., Rozet, E., Dubois, N., Charlier, C., Hubert, P., Chapelle, J.P., Krzesinski, J.M. & Delanaye, P. (2008). Performance of iohexol determination in serum and urine by HPLC: Validation, risk and uncertainty assessment, *Clin Chim Acta*,
- Chantler, C., Garnett, E.S., Parsons, V. & Veall, N. (1969). Glomerular filtration rate measurement in man by the single injection methods using 51Cr-EDTA, *Clin Sci*, Vol.37, No.1, pp. 169-180
- Cockcroft, D.W., Gault, M.H. (1976). Prediction of creatinine clearance from serum creatinine, *Nephron*, Vol.16, No.1, pp. 31-41
- Cohen, M.L., Smith, F.G., Jr., Mindell, R.S. & Vernier, R.L. (1969). A simple, reliable method of measuring glomerular filtration rate using single, low dose sodium iothalamate I-131, *Pediatrics*, Vol.43, No.3, pp. 407-415
- Corcoran, A.C. (1952). III. Determination of inulin clearance, *Methods Med Res*, Vol.5, pp. 246-248
- Dall'Amico, R., Montini, G., Pisanello, L., Piovesan, G., Bottaro, S., Cracco, A.T., Zacchello, G. & Zacchello, F. (1995). Determination of inulin in plasma and urine by reversed-phase high-performance liquid chromatography, *J Chromatogr B Biomed Appl*, Vol.672, No.1, pp. 155-159
- Dalmeida, W., Suki, W.N. (1988). Measurement of GFR with non-radioisotopic radio contrast agents, *Kidney Int*, Vol.34, No.5, pp. 725-728

- Day, D.F., Workman, W.E. (1984). A simple inulin assay for renal clearance determination using an immobilized beta-fructofuranosidase, *Ann N Y Acad Sci*, Vol.434, pp. 504-507
- De Santo, N.G., Anastasio, P., Cirillo, M., Santoro, D., Spitali, L., Mansi, L., Celentano, L., Capodicasa, D., Cirillo, E., Del, V.E., Pascale, C. & Capasso, G. (1999). Measurement of glomerular filtration rate by the 99mTc-DTPA renogram is less precise than measured and predicted creatinine clearance, *Nephron*, Vol.81, No.2, pp. 136-140
- Delanaye, P., Cohen, E.P. (2008). Formula-based estimates of the GFR: equations variable and uncertain, *Nephron Clin Pract*, Vol.110, No.1, pp. c48-c53
- Delanaye, P., Mariat, C., Maillard, N., Krzesinski, J.M. & Cavalier, E. (2011a). Are the creatinine-based equations accurate to estimate glomerular filtration rate in african american populations?, *Clin J Am Soc Nephrol*, Vol.6, No.4, pp. 906-912
- Delanaye, P., Souvignet, M., Dubourg, L., Thibaudin, L., Maillard, N., Krzesinski, J.M., Cavalier, E. & Mariat, C. (2011b). Le dosage de l'inuline: mise au point, *Ann Biol Clin (Paris)*, Vol.69, No.3, pp. 273-284
- Delanghe, J., Bellon, J., De Buyzere, M., Van Daele, G. & Leroux-Roels, G. (1991). Elimination of glucose interference in enzymatic determination of inulin, *Clin Chem*, Vol.37, No.11, pp. 2017-2018
- Denis, M.C., Venne, K., Lesiege, D., Francoeur, M., Groleau, S., Guay, M., Cusson, J. & Furtos, A. (2008). Development and evaluation of a liquid chromatography-mass spectrometry assay and its application for the assessment of renal function, *J Chromatogr A*, Vol.1189, No.1-2, pp. 410-416
- Denneberg, T., Ek, J. & Hedenskog, I. (1961). Comparison of the renal excretion of I-131-labelled hypaque and inulin, *Acta Med Scand*, Vol.170, pp. 169-181
- Ditzel, J., Vestergaard, P. & Brinklov, M. (1972). Glomerular filtration rate determined by 51 Cr-EDTA-complex. A practical method based upon the plasma disappearance curve determined from four plasma samples, *Scand J Urol Nephrol*, Vol.6, No.2, pp. 166-170
- Dodge, W.F., Travis, L.B. & Daeschner, C.W. (1967). Comparison of endogenous creatinine clearance with inulin clearance, *Am J Dis Child*, Vol.113, No.6, pp. 683-692
- Donaldson, I.M. (1968). Comparison of the renal clearances of inulin and radioactive diatrizoate ("Hypaque") as measures of the glomerular filtration rate in man, *Clin Sci*, Vol.35, No.3, pp. 513-524
- Dowling, T.C., Frye, R.F., Fraley, D.S. & Matzke, G.R. (1999). Comparison of iothalamate clearance methods for measuring GFR, *Pharmacotherapy*, Vol.19, No.8, pp. 943-950
- Dowling, T.C., Frye, R.F. & Zemaitis, M.A. (1998). Simultaneous determination of paminohippuric acid, acetyl-p-aminohippuric acid and iothalamate in human plasma and urine by high-performance liquid chromatography, *J Chromatogr B Biomed Sci Appl*, Vol.716, No.1-2, pp. 305-313
- Downes, A.M., McDonald, I.W. (1964). The chromium-51 Complex of ethylenediaminetetraacetic acid as a soluble rumen marker, *Br J Nutr*, Vol.18, pp. 153-162
- Dubourg, L., Hadj-Aissa, A. & Ferrier, B. (2010). Adaptation of an enzymatic polyfructosan assay to clinical practice, *Anal Biochem*,

- Durand, E., Chaumet-Riffaud, P., Archambaud, F., Moati, F. & Prigent, A. (2006). Mesure de la fonction rénale par les méthodes radio-isotopiques, *EMC*, Vol.Néphrologie, No.18-010-A-10, pp. 1-15
- Edelson, J., Shaw, D. & Palace, G. (1984). Pharmacokinetics of iohexol, a new nonionic radiocontrast agent, in humans, *J Pharm Sci*, Vol.73, No.7, pp. 993-995
- Effersoe, H., Rosenkilde, P., Groth, S., Jensen, L.I. & Golman, K. (1990). Measurement of renal function with iohexol. A comparison of iohexol, 99mTc-DTPA, and 51Cr-EDTA clearance, *Invest Radiol*, Vol.25, No.7, pp. 778-782
- Eide, I. (1970). Renal excretion of 51Cr-EDTA studied with stop flow technique, *Scand J Clin Lab Invest*, Vol.26, No.4, pp. 373-380
- Elwood, C.M., Sigman, E.M. (1967). The measurement of glomerular filtration rate and effective renal plasma flow in man by iothalamate 125-I and iodopyracet 131-I, *Circulation*, Vol.36, No.3, pp. 441-448
- Erley, C.M., Bader, B.D., Berger, E.D., Vochazer, A., Jorzik, J.J., Dietz, K. & Risler, T. (2001). Plasma clearance of iodine contrast media as a measure of glomerular filtration rate in critically ill patients, *Crit Care Med*, Vol.29, No.8, pp. 1544-1550
- Farthing, D., Sica, D.A., Fakhry, I., Larus, T., Ghosh, S., Farthing, C., Vranian, M. & Gehr, T. (2005). Simple HPLC-UV method for determination of iohexol, iothalamate, paminohippuric acid and n-acetyl-p-aminohippuric acid in human plasma and urine with ERPF, GFR and ERPF/GFR ratio determination using colorimetric analysis, *J Chromatogr B Analyt Technol Biomed Life Sci*, Vol.826, No.1-2, pp. 267-272
- Favre, H.R., Wing, A.J. (1968). Simultaneous 51Cr edetic acid, inulin, and endogenous creatinine clearances in 20 patients with renal disease, *Br Med J*, Vol.1, No.5584, pp. 84-86
- Fawdry, R.M., Gruenewald, S.M., Collins, L.T. & Roberts, A.J. (1985). Comparative assessment of techniques for estimation of glomerular filtration rate with 99mTc-DTPA, *Eur J Nucl Med*, Vol.11, No.1, pp. 7-12
- Fleming, J.S., Zivanovic, M.A., Blake, G.M., Burniston, M. & Cosgriff, P.S. (2004). Guidelines for the measurement of glomerular filtration rate using plasma sampling, *Nucl Med Commun*, Vol.25, No.8, pp. 759-769
- Foreman, H., Trujillo, T.T. (1954). The metabolism of C14-labeled ethylenediaminetetraacetic acid in human beings, *J Lab Clin Med*, Vol.43, pp. 566-574
- Forland, M., Pullman, T.N., Lavender, A.R. & Aho, I. (1966). The renal excretion of ethylenediaminetetraacetate in the dog, *J Pharmacol Exp Ther*, Vol.153, No.1, pp. 142-147
- Frennby, B., Sterner, G., Almen, T., Hagstam, K.E., Hultberg, B. & Jacobsson, L. (1995). The use of iohexol clearance to determine GFR in patients with severe chronic renal failure--a comparison between different clearance techniques, *Clin Nephrol*, Vol.43, No.1, pp. 35-46
- Frennby, B., Sterner, G., Almen, T., Hagstam, K.E. & Jacobsson, L. (1994). Determination of low glomerular filtration rate using iohexol clearance, *Invest Radiol*, Vol.29 Suppl 2, pp. S234-S235
- Friedman, A.N., Strother, M., Quinney, S.K., Hall, S., Perkins, S.M., Brizendine, E.J., Inman, M., Gomez, G., Shihabi, Z., Moe, S. & Li, L. (2010). Measuring the glomerular filtration rate in obese individuals without overt kidney disease, *Nephron Clin Pract*, Vol.116, No.3, pp. c224-c234

- Froissart, M., Rossert, J., Jacquot, C., Paillard, M. & Houillier, P. (2005a). Predictive performance of the modification of diet in renal disease and Cockcroft-Gault equations for estimating renal function, *J Am Soc Nephrol*, Vol.16, No.3, pp. 763-773
- Froissart, M.C., Rossert, J. & Houillier, P. (2005b). The new Mayo Clinic equation for estimating glomerular filtration rate, *Ann Intern Med*, Vol.142, No.8, pp. 679
- Gagnon, J.A., Schrier, R.W., Weis, T.P., Kokotis, W. & Mailloux, L.U. (1971). Clearance of iothalamate-125 I as a measure of glomerular filtration rate in the dog, *J Appl Physiol*, Vol.30, No.5, pp. 774-778
- Galli, G., Rufini, V., Meduri, G., Piraccini, R. & D'Andrea, G. (1994). Determination of glomerular filtration rate with 99mTc-DTPA in clinical practice, *J Nucl Biol Med*, Vol.38, No.4, pp. 556-565
- Garnett, E.S., Parsons, V. & Veall, N. (1967). Measurement of glomerular filtration-rate in man using a 51Cr-edetic-acid complex, *Lancet*, Vol.1, No.7494, pp. 818-819
- Gaspari, F., Mainardi, L., Ruggenenti, P. & Remuzzi, G. (1991). High-performance liquid chromatographic determination of iothalamic acid in human plasma and urine, *J Chromatogr*, Vol.570, No.2, pp. 435-440
- Gaspari, F., Mosconi, L., Vigano, G., Perico, N., Torre, L., Virotta, G., Bertocchi, C., Remuzzi, G. & Ruggenenti, P. (1992). Measurement of GFR with a single intravenous injection of nonradioactive iothalamate, *Kidney Int*, Vol.41, No.4, pp. 1081-1084
- Gaspari, F., Perico, N. & Remuzzi, G. (1997). Measurement of glomerular filtration rate, *Kidney Int Suppl*, Vol.63, pp. S151-S154
- Gaspari, F., Perico, N., Ruggenenti, P., Mosconi, L., Amuchastegui, C.S., Guerini, E., Daina, E. & Remuzzi, G. (1995). Plasma clearance of nonradioactive iohexol as a measure of glomerular filtration rate, *J Am Soc Nephrol*, Vol.6, No.2, pp. 257-263
- Gates, G.F. (1984). Computation of glomerular filtration rate with Tc-99m DTPA: an inhouse computer program, *J Nucl Med*, Vol.25, No.5, pp. 613-618
- Ginjaume, M., Casey, M., Barker, F. & Duffy, G. (1985). Measurement of glomerular filtration rate using technetium-99m DTPA, *J Nucl Med*, Vol.26, No.11, pp. 1347-1349
- Goates, J.J., Morton, K.A., Whooten, W.W., Greenberg, H.E., Datz, F.L., Handy, J.E., Scuderi, A.J., Haakenstad, A.O. & Lynch, R.E. (1990). Comparison of methods for calculating glomerular filtration rate: technetium-99m-DTPA scintigraphic analysis, protein-free and whole-plasma clearance of technetium-99m-DTPA and iodine-125-iothalamate clearance, *J Nucl Med*, Vol.31, No.4, pp. 424-429
- Griep, R.J., Nelp, W.B. (1969). Mechanism of excretion of radioiodinated sodium iothalamate, *Radiology*, Vol.93, No.4, pp. 807-811
- Griffiths, P.D., Drolc, Z., Green, A., Taylor, C.M. & White, R.H. (1988). Comparison of 51Cr-EDTA and 99mTc-DTPA slope clearances in children with vesicoureteric reflux, *Child Nephrol Urol*, Vol.9, No.5, pp. 283-285
- Guesry, P., Kaufman, L., Orloff, S., Nelson, J.A., Swann, S. & Holliday, M. (1975). Measurement of glomerular filtration rate by fluorescent excitation of non-radioactive meglumine iothalamate, *Clin Nephrol*, Vol.3, No.4, pp. 134-138
- Gunasekera, R.D., Allison, D.J. & Peters, A.M. (1996). Glomerular filtration rate in relation to extracellular fluid volume: similarity between 99mTc-DTPA and inulin, *Eur J Nucl Med*, Vol.23, No.1, pp. 49-54

- Gutman, Y., Gottschalk, C. & Lassiter, W.E. (1965). Micropuncture study of inulin absorption in the rat kidney, *Science*, Vol.147, pp. 753-754
- Hagstam, K.E., Nordenfelt, I., Svensson, L. & Svensson, S.E. (1974). Comparison of different methods for determination of glomerular filtration rate in renal disease, *Scand J Clin Lab Invest*, Vol.34, No.1, pp. 31-36
- Hall, P.M., Rolin, H. (1995). Iothalamate clearance and its use in large-scale clinical trials, *Curr Opin Nephrol Hypertens*, Vol.4, No.6, pp. 510-513
- Handelsman, M.B., Sass, M. (1956). The use of capillary blood for estimating renal clearance of inulin and glucose excretion by means of an anthrone procedure, *J Lab Clin Med*, Vol.48, No.5, pp. 759-768
- Harrow, B.R. (1956). Experiences in intravenous urography using hypaque, *Am J Roentgenol Radium Ther Nucl Med*, Vol.75, No.5, pp. 870-876
- Hauser, W., Atkins, H.L., Nelson, K.G. & Richards, P. (1970). Technetium-99m DTPA: a new radiopharmaceutical for brain and kidney scanning, *Radiology*, Vol.94, No.3, pp. 679-684
- Heath, D.A., Knapp, M.S. & Walker, W.H. (1968). Comparison between inulin and 51Cr-labelled edetic acid for the measurement of glomerular filtration-rate, *Lancet*, Vol.2, No.7578, pp. 1110-1112
- Hendrix, J.P., Westfall, B.B. & Richards, A.N. (1937). Quantitative studies of the composition of glomerular urine. The glomerular excretion of inulin in frogs and Necturi, *J Biol Chem*, Vol.116, No.2, pp. 735-747
- Heron, C.W., Underwood, S.R. & Dawson, P. (1984). Electrocardiographic changes during intravenous urography: a study with sodium iothalamate and iohexol, *Clin Radiol*, Vol.35, No.2, pp. 137-141
- Heyrovsky, A. (1956). A new method for determination of inulin in plasma and urine, *Clin Chim Acta*, Vol.1, No.5, pp. 470-474
- Hilson, A.J., Mistry, R.D. & Maisey, M.N. (1976). 99Tcm-DTPA for the measurement of glomerular filtration rate, *Br J Radiol*, Vol.49, No.585, pp. 794-796
- Höber, R. (1930). Beweis selektiver Sekretion durch die Tubulusepithelien der Niere, *Arch f d ges Physiol*, Vol.224, pp. 72
- Houlihan, C., Jenkins, M., Osicka, T., Scott, A., Parkin, D. & Jerums, G. (1999). A comparison of the plasma disappearance of iohexol and 99mTc-DTPA for the measurement of glomerular filtration rate (GFR) in diabetes, *Aust N Z J Med*, Vol.29, No.5, pp. 693-700
- Hubbard, R.S., Loomis, T.A. (1942). The determination of inuline, *J Biol Chem*, Vol.145, pp. 641-645
- Isaka, Y., Fujiwara, Y., Yamamoto, S., Ochi, S., Shin, S., Inoue, T., Tagawa, K., Kamada, T. & Ueda, N. (1992). Modified plasma clearance technique using nonradioactive iothalamate for measuring GFR, *Kidney Int*, Vol.42, No.4, pp. 1006-1011
- Israelit, A.H., Long, D.L., White, M.G. & Hull, A.R. (1973). Measurement of glomerular filtration rate utilizing a single subcutaneous injection of 125I-iothalamate, *Kidney Int*, Vol.4, No.5, pp. 346-349
- Jagenburg, R., Attman, P.O., Aurell, M. & Bucht, H. (1978). Determination of glomerular filtration rate in advanced renal insufficiency, Scand J Urol Nephrol, Vol.12, No.2, pp. 133-137

- Jeghers, O., Piepsz, A. & Ham, H.R. (1990). What does protein binding of radiopharmaceuticals mean exactly?, *Eur J Nucl Med*, Vol.17, No.3-4, pp. 101-102
- Johansson, R.S., Falch, D.K. (1978). 113mIn-DTPA, a useful compound for the determination of glomerular filtration rate (GFR). The binding of 113mIn to DTPA and a comparison between GFR estimated with 113mIn-DTPA and 125I-iothalamate, *Eur J Nucl Med*, Vol.3, No.3, pp. 179-181
- Jung, K., Klotzek, S. & Schulze, B.D. (1990). Refinements of assays for low concentrations of inulin in serum, *Nephron*, Vol.54, No.4, pp. 360-361
- Kainer, G., McIlveen, B., Hoschl, R. & Rosenberg, A.R. (1979). Assessment of individual renal function in children using 99mTc-DTPA, *Arch Dis Child*, Vol.54, No.12, pp. 931-936
- Kempi, V., Persson, R.B. (1975). 99mTc-DTPA(Sn) dry-kit preparation. Quality control and clearance studies, *Nucl Med (Stuttg)*, Vol.13, No.4, pp. 389-399
- Klopper, J.F., Hauser, W., Atkins, H.L., Eckelman, W.C. & Richards, P. (1972). Evaluation of 99m Tc-DTPA for the measurement of glomerular filtration rate, *J Nucl Med*, Vol.13, No.1, pp. 107-110
- Kos, T., Moser, P., Yilmatz, N., Mayer, G., Pacher, R. & Hallstrom, S. (2000). High-performance liquid chromatographic determination of p-aminohippuric acid and iothalamate in human serum and urine: comparison of two sample preparation methods, *J Chromatogr B Biomed Sci Appl*, Vol.740, No.1, pp. 81-85
- Krutzen, E., Back, S.E., Nilsson-Ehle, I. & Nilsson-Ehle, P. (1984). Plasma clearance of a new contrast agent, iohexol: a method for the assessment of glomerular filtration rate, *J Lab Clin Med*, Vol.104, No.6, pp. 955-961
- Krutzen, E., Back, S.E. & Nilsson-Ehle, P. (1990). Determination of glomerular filtration rate using iohexol clearance and capillary sampling, *Scand J Clin Lab Invest*, Vol.50, No.3, pp. 279-283
- Kuehnle, H.F., von Dahl, K. & Schmidt, F.H. (1992). Fully enzymatic inulin determination in small volume samples without deproteinization, *Nephron*, Vol.62, No.1, pp. 104-107
- Kwong, Y.T., Stevens, L.A., Selvin, E., Zhang, Y.L., Greene, T., Van, L.F., Levey, A.S. & Coresh, J. (2010). Imprecision of urinary iothalamate clearance as a gold-standard measure of GFR decreases the diagnostic accuracy of kidney function estimating equations, *Am J Kidney Dis*, Vol.56, No.1, pp. 39-49
- Laake, H. (1954). Inulin clearance studies; concerning the cause of the reduced clearance figures in successive periods after one injection of inulin, *Acta Med Scand*, Vol.148, No.2, pp. 135-146
- Ladegaard-Pedersen, H.J. (1972). Measurement of extracellular volume and renal clearance by a single injection of inulin, *Scand J Clin Lab Invest*, Vol.29, No.2, pp. 145-153
- LaFrance, N.D., Drew, H.H. & Walser, M. (1988). Radioisotopic measurement of glomerular filtration rate in severe chronic renal failure, *J Nucl Med*, Vol.29, No.12, pp. 1927-1930
- Lavender, S., Hilton, P.J. & Jones, N.F. (1969). The measurement of glomerular filtration-rate in renal disease, *Lancet*, Vol.2, No.7632, pp. 1216-1218
- Lee, S.Y., Chun, M.R., Kim, D.J. & Kim, J.W. (2006). Determination of iohexol clearance by high-performance liquid chromatography-tandem mass spectrometry (HPLC-MS/MS), *J Chromatogr B Analyt Technol Biomed Life Sci*, Vol.839, No.1-2, pp. 124-129

- Lentjes, E.G., Florijn, K.W., Chang, P.C. & van Dam, W. (1994). Inulin measurement in serum and urine with an autoanalyser, corrected for glucose interference, *Eur J Clin Chem Clin Biochem*, Vol.32, No.8, pp. 625-628
- Levey, A.S., Bosch, J.P., Lewis, J.B., Greene, T., Rogers, N. & Roth, D. (1999). A more accurate method to estimate glomerular filtration rate from serum creatinine: a new prediction equation. Modification of Diet in Renal Disease Study Group, *Ann Intern Med*, Vol.130, No.6, pp. 461-470
- Levey, A.S., Coresh, J., Greene, T., Stevens, L.A., Zhang, Y.L., Hendriksen, S., Kusek, J.W. & Van Lente, F. (2006). Using standardized serum creatinine values in the modification of diet in renal disease study equation for estimating glomerular filtration rate, *Ann Intern Med*, Vol.145, No.4, pp. 247-254
- Levey AS, Madaio MP, Perrone RD 1991: Laboratory assessment of renal disease: clearance, urinalysis, and renal biopsy. In: Brenner BM, Rector FC (eds): The kidney. W.B. Saunders Company, Philadelphia, pp. 919-968
- Levey, A.S., Stevens, L.A., Schmid, C.H., Zhang, Y.L., Castro, A.F., III, Feldman, H.I., Kusek, J.W., Eggers, P., Van Lente, F., Greene, T. & Coresh, J. (2009). A new equation to estimate glomerular filtration rate, *Ann Intern Med*, Vol.150, No.9, pp. 604-612
- Lewis, R., Kerr, N., Van Buren, C., Lowry, P., Sandler, C., Frazier, O.H., Powers, P., Herson, J., Corriere, J., Jr., Kerman, R. & . (1989). Comparative evaluation of urographic contrast media, inulin, and 99mTc-DTPA clearance methods for determination of glomerular filtration rate in clinical transplantation, *Transplantation*, Vol.48, No.5, pp. 790-796
- Lingardh, G. (1972). Renal clearance investigations with 51 Cr-EDTA and 125 I-hippuran, *Scand J Urol Nephrol*, Vol.6, No.1, pp. 63-71
- Little, J.M. (1949). A modified diphenylamine procedure for the determination of inulin, *J Biol Chem*, Vol.180, No.2, pp. 747-754
- Lundqvist, S., Hietala, S.O., Berglund, C. & Karp, K. (1994). Simultaneous urography and determination of glomerular filtration rate. A comparison of total plasma clearances of iohexol and 51Cr-EDTA in plegic patients, *Acta Radiol*, Vol.35, No.4, pp. 391-395
- Lundqvist, S., Hietala, S.O., Groth, S. & Sjodin, J.G. (1997). Evaluation of single sample clearance calculations in 902 patients. A comparison of multiple and single sample techniques, *Acta Radiol*, Vol.38, No.1, pp. 68-72
- Ma, Y.C., Zuo, L., Zhang, C.L., Wang, M., Wang, R.F. & Wang, H.Y. (2007). Comparison of 99mTc-DTPA renal dynamic imaging with modified MDRD equation for glomerular filtration rate estimation in Chinese patients in different stages of chronic kidney disease, *Nephrol Dial Transplant*, Vol.22, No.2, pp. 417-423
- Mafham, M.M., Niculescu-Duvaz, I., Barron, J., Emberson, J.R., Dockrell, M.E., Landray, M.J. & Baigent, C. (2007). A practical method of measuring glomerular filtration rate by iohexol clearance using dried capillary blood spots, *Nephron Clin Pract*, Vol.106, No.3, pp. c104-c112
- Maher, F.T., Nolan, N.G. & Elveback, L.R. (1971). Comparison of simultaneous clearances of 125-I-labeled sodium lothalamate (Glofil) and of inulin, *Mayo Clin Proc*, Vol.46, No.10, pp. 690-691
- Maher, F.T., Tauxe, W.N. (1969). Renal clearance in man of pharmaceuticals containing radioactive iodine, *JAMA*, Vol.207, No.1, pp. 97-104

- Malamos, B., Dontas, A.S., Koutras, D.A., Marketos, S., Sfontouris, J. & Papanicolaou, N. (1967). 125I-sodium iothalamate in the determination of the glomerular filtration rate, *Nucl Med (Stuttg)*, Vol.6, No.3, pp. 304-310
- McChesney, E.W., Hoppe, J.O. (1957). Studies of the tissue distribution and excretion of sodium diatrizoate in laboratory animals, *Am J Roentgenol Radium Ther Nucl Med*, Vol.78, No.1, pp. 137-144
- Medeiros, F.S., Sapienza, M.T., Prado, E.S., Agena, F., Shimizu, M.H., Lemos, F.B., Buchpiguel, C.A., Ianhez, L.E. & David-Neto, E. (2009). Validation of plasma clearance of 51Cr-EDTA in adult renal transplant recipients: comparison with inulin renal clearance, *Transpl Int*, Vol.22, No.3, pp. 323-331
- Meschan, I., Deyton, W.E., Schmid, H.E. & Watts, F.C. (1963). The utilization of I 131-labeled renografin as an inulin substitute for renal clearance rate determination, *Radiology*, Vol.81, pp. 974-979
- Mirouze, J., Barjon, P., Monnier, L. & Mimran, A. (1972). Détermination de la filtration glomérulaire par l'iothalamate radioactif: comparaison des méthodes par perfusion continue et injection unique rapide, *Vie Med Can Fr*, Vol.1, No.9, pp. 837-841
- Möller, E., McIntosh, J.F. & Van Slycke, D.D. (1929). Studies of urea excretion. II. Relationship between urine volume and the rate of urea excretion by normal adults, *J Clin Invest*, Vol.6, No.3, pp. 427-465
- Morgan, D.B., Dillon, S. & Payne, R.B. (1978). The assessment of glomerular function: creatinine clearance or plasma creatinine?, *Postgrad Med J*, Vol.54, No.631, pp. 302-310
- Morris, A.M., Elwood, C., Sigman, E.M. & Catanzaro, A. (1965). The renal clearance of 131-I labeled meglumine diatrizoate (renografin) in man, *J Nucl Med*, Vol.6, pp. 183-191
- Mulligan, J.S., Blue, P.W. & Hasbargen, J.A. (1990). Methods for measuring GFR with technetium-99m-DTPA: an analysis of several common methods, *J Nucl Med*, Vol.31, No.7, pp. 1211-1219
- Mutzel, W., Siefert, H.M. & Speck, U. (1980). Biochemical-pharmacologic properties of iohexol, *Acta Radiol Suppl*, Vol.362, pp. 111-115
- Niculescu-Duvaz, I., D'Mello, L., Maan, Z., Barron, J.L., Newman, D.J., Dockrell, M.E. & Kwan, J.T. (2006). Development of an outpatient finger-prick glomerular filtration rate procedure suitable for epidemiological studies, *Kidney Int*, Vol.69, No.7, pp. 1272-1275
- Nilsson-Ehle, P. (2002). Iohexol clearance for the determination of glomerular filtration rate: 15 years' experience in clinical practice, *eJIFCC*, Vol.13, No.2,
- Nilsson-Ehle, P., Grubb, A. (1994). New markers for the determination of GFR: iohexol clearance and cystatin C serum concentration, *Kidney Int Suppl*, Vol.47, pp. S17-S19
- Nossen, J.O., Jakobsen, J.A., Kjaersgaard, P., Andrew, E., Jacobsen, P.B. & Berg, K.J. (1995). Elimination of the non-ionic X-ray contrast media iodixanol and iohexol in patients with severely impaired renal function, *Scand J Clin Lab Invest*, Vol.55, No.4, pp. 341-350
- Nosslin, B. (1965). Determination of clearance and distribution volume with the single injection method, *Acta Med Scand Suppl*, Vol.442, pp. 97-101
- Notghi, A., Merrick, M.V., Ferrington, C. & Anderton, J.L. (1986). A comparison of simplified and standard methods for the measurement of glomerular filtration rate and renal tubular function, *Br J Radiol*, Vol.59, No.697, pp. 35-39

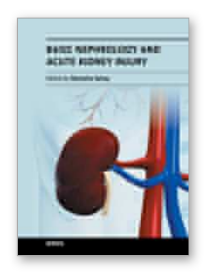
- O'Reilly, P.H., Brooman, P.J., Martin, P.J., Pollard, A.J., Farah, N.B. & Mason, G.C. (1986). Accuracy and reproducibility of a new contrast clearance method for the determination of glomerular filtration rate, *Br Med J (Clin Res Ed)*, Vol.293, No.6541, pp. 234-236
- O'Reilly, P.H., Jones, D.A. & Farah, N.B. (1988). Measurement of the plasma clearance of urographic contrast media for the determination of glomerular filtration rate, *J Urol*, Vol.139, No.1, pp. 9-11
- Odlind, B., Hallgren, R., Sohtell, M. & Lindstrom, B. (1985). Is 125I iothalamate an ideal marker for glomerular filtration?, *Kidney Int*, Vol.27, No.1, pp. 9-16
- Olofsson, P., Krutzen, E. & Nilsson-Ehle, P. (1996). Iohexol clearance for assessment of glomerular filtration rate in diabetic pregnancy, Eur J Obstet Gynecol Reprod Biol, Vol.64, No.1, pp. 63-67
- Olsson, B., Aulie, A., Sveen, K. & Andrew, E. (1983). Human pharmacokinetics of iohexol. A new nonionic contrast medium, *Invest Radiol*, Vol.18, No.2, pp. 177-182
- Ott, N.T. (1975). A simple technique for estimating glomerular filtration rate with subcutaneous injection of (125I)lothalamate, *Mayo Clin Proc*, Vol.50, No.11, pp. 664-668
- Owen, J.E., Walker, R.G., Willems, D., Guignard, P.A. & d'Apice, A.J. (1982). Cadmium telluride detectors in the external measurement of glomerular filtration rate using 99mTc-DTPA (Sn): comparison with 51Cr-EDTA and 99mTc-DTPA (Sn) plasma sample methods, *Clin Nephrol*, Vol.18, No.4, pp. 200-203
- Owman, T., Olin, T. (1978). Biliary excretion of urographic contrast media (iothalamate, diatrizoate, P 286 and metrizamide), *Ann Radiol (Paris)*, Vol.21, No.4-5, pp. 309-314
- Pastore, A., Bernardini, S., Dello, S.L., Rizzoni, G., Cortese, C. & Federici, G. (2001). Simultaneous determination of inulin and p-aminohippuric acid in plasma and urine by reversed-phase high-performance liquid chromatography, *J Chromatogr B Biomed Sci Appl*, Vol.751, No.1, pp. 187-191
- Perrone, R.D., Steinman, T.I., Beck, G.J., Skibinski, C.I., Royal, H.D., Lawlor, M. & Hunsicker, L.G. (1990). Utility of radioisotopic filtration markers in chronic renal insufficiency: simultaneous comparison of 125I-iothalamate, 169Yb-DTPA, 99mTc-DTPA, and inulin. The Modification of Diet in Renal Disease Study, *Am J Kidney Dis*, Vol.16, No.3, pp. 224-235
- Petri, M., Bockenstedt, L., Colman, J., Whiting-O'Keefe, Q., Fitz, G., Sebastian, A. & Hellmann, D. (1988). Serial assessment of glomerular filtration rate in lupus nephropathy, *Kidney Int*, Vol.34, No.6, pp. 832-839
- Piepsz, A., Erbsmann, F., Vainsel, M. & Cantraine, F. (1974). Determination of glomerular filtration rate in children by external measurement of the disappearance of 113m-indium-DTPA in plasma, *Biomedicine*, Vol.20, No.3, pp. 192-197
- Prueksaritanont, T., Chen, M.L. & Chiou, W.L. (1984). Simple and micro high-performance liquid chromatographic method for simultaneous determination of paminohippuric acid and iothalamate in biological fluids, *J Chromatogr*, Vol.306, pp. 89-97
- Prueksaritanont, T., Lui, C.Y., Lee, M.G. & Chiou, W.L. (1986). Renal and non-renal clearances of iothalamate, *Biopharm Drug Dispos*, Vol.7, No.4, pp. 347-355

- Pucci, L., Bandinelli, S., Penno, G., Nannipieri, M., Rizzo, L. & Navalesi, R. (1998). Iohexol plasma clearance in determining glomerular filtration rate in diabetic patients, *Ren Fail*, Vol.20, No.2, pp. 277-284
- Pucci, L., Bandinelli, S., Pilo, M., Nannipieri, M., Navalesi, R. & Penno, G. (2001). Iohexol as a marker of glomerular filtration rate in patients with diabetes: comparison of multiple and simplified sampling protocols, *Diabet Med*, Vol.18, No.2, pp. 116-120
- Reba, R.C., Hosain, F. & Wagner, H.N., Jr. (1968). Indium-113m diethylenetriaminepentaacetic acid (DTPA): a new radiopharmaceutical for study of the kidneys, *Radiology*, Vol.90, No.1, pp. 147-149
- Rehberg, P.B. (1926a). Studies on Kidney Function: The Excretion of urea and chlorine analysed according to a modified filtration-reabsorption theory, *Biochem J*, Vol.20, No.3, pp. 461-482
- Rehberg, P.B. (1926b). Studies on Kidney Function: The Rate of Filtration and Reabsorption in the Human Kidney, *Biochem J*, Vol.20, No.3, pp. 447-460
- Rehling, M., Moller, M.L., Thamdrup, B., Lund, J.O. & Trap-Jensen, J. (1984). Simultaneous measurement of renal clearance and plasma clearance of 99mTc-labelled diethylenetriaminepenta-acetate, 51Cr-labelled ethylenediaminetetra-acetate and inulin in man, *Clin Sci (Lond)*, Vol.66, No.5, pp. 613-619
- Rehling, M., Moller, M.L., Thamdrup, B., Lund, J.O. & Trap-Jensen, J. (1986). Reliability of a 99mTc-DTPA gamma camera technique for determination of single kidney glomerular filtration rate. A comparison to plasma clearance of 51Cr-EDTA in one-kidney patients, using the renal clearance of inulin as a reference, *Scand J Urol Nephrol*, Vol.20, No.1, pp. 57-62
- Rehling, M., Nielsen, B.V., Pedersen, E.B., Nielsen, L.E., Hansen, H.E. & Bacher, T. (1995). Renal and extrarenal clearance of 99mTc-MAG3: a comparison with 125I-OIH and 51Cr-EDTA in patients representing all levels of glomerular filtration rate, *Eur J Nucl Med*, Vol.22, No.12, pp. 1379-1384
- Rehling, M., Nielsen, L.E. & Marqversen, J. (2001). Protein binding of 99Tcm-DTPA compared with other GFR tracers, *Nucl Med Commun*, Vol.22, No.6, pp. 617-623
- Reidenberg, M.M., Lorenzo, B.J., Drayer, D.E., Kluger, J., Nestor, T., Regnier, J.C., Kowal, B.A. & Bekersky, I. (1988). A nonradioactive iothalamate method for measuring glomerular filtration rate and its use to study the renal handling of cibenzoline, *Ther Drug Monit*, Vol.10, No.4, pp. 434-437
- Richards, A.N., Westfall, B.B. & Bott, P.A. (1934). Renal excretion of inulin, creatinine, and xylose in normal dogs., *Proc Soc Exp Biol Med*, Vol.32, pp. 73
- Robson, J.S., Ferguson, M.H., Olbrich, O. & Stewart, C.P. (1949). The determination of the renal clearance of inulin in man, *Q J Exp Physiol*, Vol.35, No.2, pp. 111-134
- Rodby, R.A., Ali, A., Rohde, R.D. & Lewis, E.J. (1992). Renal scanning 99mTc diethylene-triamine pentaacetic acid glomerular filtration rate (GFR) determination compared with iothalamate clearance GFR in diabetics. The Collaborative Study Group for The study of Angiotensin-Converting Enzyme Inhibition in Diabetic Nephropathy, *Am J Kidney Dis*, Vol.20, No.6, pp. 569-573
- Roe, J.H. (1934). A colorimetric method for the determination of fructose in blood and urine, *J Biol Chem*, Vol.107, No.1, pp. 15-21
- Rolf, D., Surtshin, A. & White, H.L. (1949). A modified diphenylamine procedure for fructose or inulin determination, *Proc Soc Exp Biol Med*, Vol.72, No.2, pp. 351-354

- Rootwelt, K., Falch, D. & Sjokvist, R. (1980). Determination of glomerular filtration rate (GFR) by analysis of capillary blood after single shot injection of 99mTc-DTPA. A comparison with simultaneous 125I-iothalamate GFR estimation showing equal GFR but difference in distribution volume, *Eur J Nucl Med*, Vol.5, No.2, pp. 97-102
- Rosenbaum, R.W., Hruska, K.A., Anderson, C., Robson, A.M., Slatopolsky, E. & Klahr, S. (1979). Inulin: an inadequate marker of glomerular filtration rate in kidney donors and transplant recipients?, *Kidney Int*, Vol.16, No.2, pp. 179-186
- Ruo, T.I., Wang, Z., Dordal, M.S. & Atkinson, A.J., Jr. (1991). Assay of inulin in biological fluids by high-performance liquid chromatography with pulsed amperometric detection, *Clin Chim Acta*, Vol.204, No.1-3, pp. 217-222
- Russell, C.D. (1987). Estimation of glomerular filtration rate using 99mTc-DTPA and the gamma camera, *Eur J Nucl Med*, Vol.12, No.11, pp. 548-552
- Russell, C.D., Bischoff, P.G., Kontzen, F., Rowell, K.L., Yester, M.V., Lloyd, L.K., Tauxe, W.N. & Dubovsky, E.V. (1985). Measurement of glomerular filtration rate using 99mTc-DTPA and the gamma camera: a comparison of methods, *Eur J Nucl Med*, Vol.10, No.11-12, pp. 519-521
- Russell, C.D., Bischoff, P.G., Rowell, K.L., Kontzen, F., Lloyd, L.K., Tauxe, W.N. & Dubovsky, E.V. (1983). Quality control of Tc-99m DTPA for measurement of glomerular filtration: concise communication, *J Nucl Med*, Vol.24, No.8, pp. 722-727
- Sambataro, M., Thomaseth, K., Pacini, G., Robaudo, C., Carraro, A., Bruseghin, M., Brocco, E., Abaterusso, C., DeFerrari, G., Fioretto, P., Maioli, M., Tonolo, G.C., Crepaldi, G. & Nosadini, R. (1996). Plasma clearance rate of 51Cr-EDTA provides a precise and convenient technique for measurement of glomerular filtration rate in diabetic humans, *J Am Soc Nephrol*, Vol.7, No.1, pp. 118-127
- Schanker, L.S., Hogben, C.A. (1961). Biliary excretion of inulin, sucrose, and mannitol: analysis of bile formation, *Am J Physiol*, Vol.200, pp. 1087-1090
- Schwartz, G.J., Furth, S., Cole, S.R., Warady, B. & Munoz, A. (2006). Glomerular filtration rate via plasma iohexol disappearance: pilot study for chronic kidney disease in children, *Kidney Int*, Vol.69, No.11, pp. 2070-2077
- Seegmiller, J.C., Burns, B.E., Fauq, A.H., Mukhtar, N., Lieske, J.C. & Larson, T.S. (2010). Iothalamate quantification by tandem mass spectrometry to measure glomerular filtration rate, *Clin Chem*, Vol.56, No.4, pp. 568-574
- Shaffer, P.A., Somogoyi, M. (1933). Copper-iodometric reagents for sugar determination, *J Biol Chem*, Vol.100, pp. 695-713
- Shannon, J.A. (1934). The excretion of inulin by the dogfish, *Squalus acanthias*, *J Cell and Comp Physiol*, Vol.5, No.3, pp. 301-310
- Shannon, J.A., Smith, H.W. (1935). The excretion of inulin, xylose, and urea by normal and phorizinized man, *J Clin Invest*, Vol.14, No.4, pp. 393-401
- Sharma, A.K., Mills, M.S., Grey, V.L. & Drummond, K.N. (1997). Infusion clearance of subcutaneous iothalamate versus standard renal clearance, *Pediatr Nephrol*, Vol.11, No.6, pp. 711-713
- Shemesh, O., Golbetz, H., Kriss, J.P. & Myers, B.D. (1985). Limitations of creatinine as a filtration marker in glomerulopathic patients, *Kidney Int*, Vol.28, No.5, pp. 830-838
- Shihabi, Z.K., Constantinescu, M.S. (1992). Iohexol in serum determined by capillary electrophoresis, *Clin Chem*, Vol.38, No.10, pp. 2117-2120

- Sigman, E.M., Elwood, C., Reagan, M.E., MorrisS, A.M. & Catanzaro, A. (1965a). The renal clearance of of I-131 labelled iothalamate in man, *Invest Urol*, Vol.2, pp. 432-438
- Sigman, E.M., Elwood, C.M. & Knox, F. (1965b). The measurement of glomerular filtration rate in man with sodium iothalamate 131-I (Conray), *J Nucl Med*, Vol.7, No.1, pp. 60-68
- Skluzacek, P.A., Szewc, R.G., Nolan, C.R., III, Riley, D.J., Lee, S. & Pergola, P.E. (2003). Prediction of GFR in liver transplant candidates, *Am J Kidney Dis*, Vol.42, No.6, pp. 1169-1176
- Skov, P.E. (1970). Glomerular filtration rate in patients with severe and very severe renal insufficiency. Determined by simultaneous inulin, creatinine and 125 iothalamate clearance, *Acta Med Scand*, Vol.187, No.5, pp. 419-428
- Smith HW 1951a: Measurement of the filtration rate. The kidney: Structure and function in health and disease. Oxford University Press Inc, New York, pp. 39-62
- Smith HW 1951b The kidney: Structure and function in health and disease. Oxford University Press Inc, New York
- Smith HW 1951c: The reliability of inulin as a measure of glomerular filtration. The kidney: Structure and function in health and disease. Oxford University Press Inc, New York, pp. 231-238
- Spencer, H., Samachson, J. & Laszlo, D. (1958). Effects of ethylenediaminetetraacetic acid on radiostrontium excretion in man, *Proc Soc Exp Biol Med*, Vol.97, pp. 565-567
- Stacy, B.D., Thorburn, G.D. (1966). Chromium-51 ethylenediaminetetraacetate for estimation of globerular filtration rate, *Science*, Vol.152, No.725, pp. 1076-1077
- Stamp, T.C., Stacey, T.E. & Rose, G.A. (1970). Comparison of glomerular filtration rate measurements using inulin, 51CrEDTA, and a phosphate infusion technique, *Clin Chim Acta*, Vol.30, No.2, pp. 351-358
- Steinitz, K. (1938). A colorimetric method for the determination of inulin in blood plasma and urine, *J Biol Chem*, Vol.126, No.2, pp. 589-593
- Sterner, G., Frennby, B., Mansson, S., Nyman, U., van Westen, D. & Almen, T. (2008). Determining 'true' glomerular filtration rate in healthy adults using infusion of inulin and comparing it with values obtained using other clearance techniques or prediction equations, *Scand J Urol Nephrol*, Vol.42, No.3, pp. 278-285
- Stevens, L.A., Levey, A.S. (2009). Measured GFR as a confirmatory test for estimated GFR, *J Am Soc Nephrol*, Vol.20, No.11, pp. 2305-2313
- Stokes, J.M., Conklin, J.W. & Huntley, H.C. (1962). Measurement of glomerular filtration rate by contrast media containing I-131 isotopes, *J Urol*, Vol.87, pp. 630-633
- Stolz, A., Hoizey, G., Toupance, O., Lavaud, S., Vitry, F., Chanard, J. & Rieu, P. (2010). Evaluation of sample bias for measuring plasma iohexol clearance in kidney transplantation, *Transplantation*, Vol.89, No.4, pp. 440-445
- Summerfield, A.L., Hortin, G.L., Smith, C.H., Wilhite, T.R. & Landt, M. (1993). Automated enzymatic analysis of inulin, *Clin Chem*, Vol.39, No.11 Pt 1, pp. 2333-2337
- Tepe, P.G., Tauxe, W.N., Bagchi, A., Rezende, P. & Krishnaiah, P.R. (1987). Comparison of measurement of glomerular filtration rate by single sample, plasma disappearance slope/intercept and other methods, *Eur J Nucl Med*, Vol.13, No.1, pp. 28-31
- Tessitore, N., Lo, S.C., Corgnati, A., Previato, G., Valvo, E., Lupo, A., Chiaramonte, S., Messa, P., D'Angelo, A., Zatti, M. & Maschio, G. (1979). 125I-iothalamate and

- creatinine clearances in patients with chronic renal diseases, *Nephron*, Vol.24, No.1, pp. 41-45
- Thomsen, H.S., Hvid-Jacobsen, K. (1991). Estimation of glomerular filtration rate from low-dose injection of iohexol and a single blood sample, *Invest Radiol*, Vol.26, No.4, pp. 332-336
- van de Wiele C., van den Eeckhaut A., Verweire, W., van Haelst, J.P., Versijpt, J. & Dierckx, R.A. (1999). Absolute 24 h quantification of 99Tcm-DMSA uptake in patients with severely reduced kidney function: a comparison with 51Cr-EDTA clearance, *Nucl Med Commun*, Vol.20, No.9, pp. 829-832
- Visser, F.W., Muntinga, J.H., Dierckx, R.A. & Navis, G. (2008). Feasibility and impact of the measurement of extracellular fluid volume simultaneous with GFR by 125I-iothalamate, *Clin J Am Soc Nephrol*, Vol.3, No.5, pp. 1308-1315
- Weber, A.F., Lee, D.W., Opheim, K. & Smith, A.L. (1985). Quantitation of iothalamate in serum and urine by high-performance liquid chromatography, *J Chromatogr*, Vol.337, No.2, pp. 434-440
- Welling, P.G., Mosegaard, A., Dobrinska, M.R. & Madsen, P.O. (1976). Pharmacokinetics of 125I-iothalamate and 131I-o-iodohippurate in man, *J Clin Pharmacol*, Vol.16, No.2-3, pp. 142-148
- Wharton, W.W., III, Sondeen, J.L., McBiles, M., Gradwohl, S.E., Wade, C.E., Ciceri, D.P., Lehmann, H.G., Stotler, R.E., Henderson, T.R., Whitaker, W.R. & . (1992). Measurement of glomerular filtration rate in ICU patients using 99mTc-DTPA and inulin, *Kidney Int*, Vol.42, No.1, pp. 174-178
- Wilson, D.M., Bergert, J.H., Larson, T.S. & Liedtke, R.R. (1997). GFR determined by nonradiolabeled iothalamate using capillary electrophoresis, *Am J Kidney Dis*, Vol.30, No.5, pp. 646-652
- Winter, C.C., Taplin, G.V. (1958). A clinical comparison and analysis of radioactive diodarast, hypaque, miokon and urokon renograms as tests of kidney function, *J Urol*, Vol.79, No.3, pp. 573-579
- Winterborn, M.H., Beetham, R. & White, R.H. (1977). Comparison of plasma disappearance and standard clearance techniques for measuring glomerular filtration rate in children with and without vesico-ureteric reflux, *Clin Nephrol*, Vol.7, No.6, pp. 262-270
- Woodruff, M.W., Malvin, R.L. (1960). Localization of renal contrast media excretion by stop flow analysis, *J Urol*, Vol.84, pp. 677-684



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