I would like to thank you for reading this information pack about Wolfram Syndrome. Wolfram Syndrome is a rare progressive neuro degenerative condition with limited life expectancy and no cure. It affects 1 in 770,000 in the UK. There are currently about 90 people (adults and children) diagnosed with WS and it is thought that there are at least another 50 people either undiagnosed or misdiagnosed.

The aim of this pack is to inform medical professionals that may see a child or adult with WS, what to be aware of to aid a quicker diagnosis and so prevent years of uncomfortable and intrusive testing. Please share the information amongst the staff in your department.

Inside this pack you will find the following:

Information sheets regarding:

Urology
Neurology
Ophthalmology
Paediatricians and Healthcare Professionals
Endocrinologists
Diabetologists
Wolfram Syndrome MDT Clinical Guidelines
WSUK Charity leaflet
Letter from Dr Fumi Urano

If you would like any further information about Wolfram Syndrome then please contact the office either by phone or email.

Yours sincerely

Tracy

Tracy Lynch
Chief Executive & Co-Founder

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Reg Charity No: 1152445. Reg Charity in England and Wales
Consider Wolfram Syndrome

Does a child under the age of 16 years complain of:

- Decreasing visual acuity / ‘greying of vision’
- Loss of colour vision

Does the child have any of these diagnostic features?

- Progressive Optic Atrophy
- Diabetes Mellitus

Does the child have any supportive features?

- Diabetes insipidus
- Sensorineural deafness
- Neurological signs (ataxia, epilepsy, neuropathy, cognitive impairment)
- Renal tract abnormalities
- Family history of Wolfram Syndrome

Look for additional features

- Cataracts
- Nystagmus
- Poor papillary reflexes
- Hypogonadism (males)
- Psychiatric disorders
- Gastrointestinal disorders

Investigations:

- Visual fields - Central with peripheral scotomas
- VEP - normal latency but low amplitude
- Genetic tests WFS1 / CISD2
- OCT - RNFL analysis
- ERG
- Absence of type 1 diabetes auto-antibodies
- MRI generalised brain atrophy with reduced signal from optic nerve and chiasm

Management:

- Correct refractive error
- Rehabilitation
- Vision support: Magnifying glasses, digital systems, voice systems
- Educational support: Touch screen laptops, mobile devices, networking
- Filtration glasses if photosensitive

Further information Wolfram Multidisciplinary Services

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University Hospitals Birmingham NHS Foundation Trust

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Version2 August 2014 - Miss Archana Kuikarni, Mr. John Ainsworth
Wolfram syndrome guide for diabetologists

The combination of insulin dependent diabetes mellitus presenting under 15 years and progressive optic atrophy is pathognomonic for Wolfram syndrome. The positive predictive value and negative predictive values are estimated at 85% (Barrett et al 1995).

Diagnostic criteria for diabetes mellitus are based on plasma blood glucose measurements and the presence or absence of symptoms. Diabetes is diagnosed when:

- A fasting plasma glucose (FPG) is ≥ 7.0mmol/L (126mg/dl) (on two occasions if there are no symptoms of diabetes).
- OR the post challenge plasma glucose is >11.1mmol/L (200mg/dl) 2 hours after a glucose load containing the equivalent of 1.75mg/kg (max 75g) of anhydrous glucose dissolved in water
- OR there are symptoms of diabetes and a random plasma glucose ≥11.1mmol/L (300mg/dl). The symptoms may include polyuria, polydipsia, and unexplained weight loss.

The diagnosis of diabetes mellitus is usually confirmed quickly in symptomatic children by measurement of an elevated blood glucose level. In this situation, if ketones are present in the blood or urine, treatment is urgent.

Type 1 diabetes associated auto-antibodies are most often absent (glutamate decarboxylase (GAD), tyrosine phosphatase (IA-2) and insulin antibodies, if available islet cell Ab (ICA) or ZnT8 Ab). Absolute insulin deficiency does occur, with ketosis, but insulopaenia is common, and may be assessed by basal and/or post stimulated C-Peptide measurements.

Management of diabetes mellitus

**Intensive education**

The principles are similar to management of type 1 diabetes. Intensive education is needed regarding insulin injection techniques, dosage adjustment, blood glucose and ketone testing, exercise, nutrition, formal smoking prevention and cessation, prevention and management of DKA and hypoglycaemia.

**Glycemic targets**

The aim is to improve metabolic control to reduce diabetes-related complications with strategies tailored to each individual, according to individual risk factors and vulnerability to severe hypoglycemia. HbA1c goals should be adapted to age: in non pregnant adult, <7%, for a child between 6 and 12yr, 7.5% and for an adolescent (12-19yr) <7.5%. Less stringent A1C goals may be appropriate for patients with a history of severe hypoglycemia, limited life expectancy, advanced microvascular or macrovascular complications, and/or extensive comorbid conditions.

**Insulin therapy**

Insulin regimen chosen according to age, duration of diabetes, lifestyle, socioeconomic factors, and family, patient and physician preferences. Intensive management is frequently required: continuous subcutaneous insulin infusion or multiple daily injection regimens using basal insulin analogues. Use insulin pen with audible signal of insulin dose delivery if available.

**Glucose monitoring**

Self-monitoring of blood glucose (adapted devices for blind people) and quarterly HbA1c measurement. If necessary and available, Continuous Glucose Monitoring System (CGMS) can be used.

**Nutrition**

Regular evaluation (at least annually) with dietetic advice (based on the nutritional needs, eating habits, lifestyle, ability and interest) ensuring normal growth and development without disturbing glycemia.

**Hypoglycemia**

Significant risk of hypoglycemia often necessitates less stringent glycemic goals or the use of a continuous glucose monitoring system. Severe hypoglycemia should be treated with intravenous dextrose (hospital) or subcutaneous glucagon (at home) according to local protocols for type 1 diabetes.

Management of diabetes complications
**Nephropathy**
- Yearly screening, starting at 12 years of age, or in patients with duration of diabetes >5 years
  - First morning urine albumin to creatinine ratio, and persistence of elevation demonstrated.

**Retinopathy**
- Yearly screening in patients with duration of diabetes more than 5 years
  - Fundoscopy, OCT scan and fluorescein angiography if signs of diabetic retinopathy are present

**Neuropathy**
- Yearly neurological exam to look for numbness, pain, cramps and paresthesia
  - Nerve conduction studies and dysautonomia assessment in presence of clinical signs or symptoms

**Dyslipidemia**
- Screen at 12 and 17y (when stabilized), or <12y if risk factors exist (obesity, familial hyperlipidemia...)
  - Fasting total cholesterol, high-density and low-density lipoprotein cholesterol, triglycerides

**Hypertension**
- Screen at least twice a year, use appropriate cuff size, +/- 24 hour ambulatory blood pressure monitoring
  - Lifestyle modification and anti-hypertensive drug therapy
Wolfram syndrome guide for endocrinologists

Apart from diabetes mellitus, other common endocrine findings in Wolfram syndrome include:

**Diabetes insipidus.**
Diabetes insipidus of central origin occurred in 72% with a median age of onset of 15.5 years (Barrett et al 1995). The range in age of onset is broad, possibly because of delays in establishing the correct diagnosis. Common symptoms include polyuria and polydipsia; the differential diagnosis includes polyuria secondary to poor glycemic control, and neuropathic bladder.

Useful investigations include 24 hour urine collection to assess volume, particularly if the patient denies symptoms. To make the diagnosis of cranial diabetes insipidus, an assessment of the concentrating ability of the urine is required. It is easiest to collect morning paired fasting urine and fasting plasma for osmolality and sodium concentration. Water deprivation tests are best avoided as they can be dangerous. A urine osmolality > 500mOsmol/L with normal serum sodium (up to 145mmol/L) and serum osmolarity (up to 295mOsmol/L) in the presence of normal serum glucose effectively excludes diabetes insipidus. A confirmation of diabetes insipidus would by a urine osmolality <150mOsmol/L, with serum Na > 145, and serum osmo > 295mOsmol/L.

Management is with desmopressin replacement according to local practices. The options are usually intranasal, buccal or oral. The intranasal preparations are about 20 times more potent than the oral, and about 15 times more potent than the buccal preparations. A safe starting dose in a child over 5 years would be 2.5 micrograms intranasal at night; and for an adult, 5-10 micrograms intranasal. The dose needs to be titrated according to symptoms, and by blood and urine biochemistry.

**Hypogonadism.**
Hypogonadism is more prevalent in males than in females. It can be either hypogonadotrophic (i.e., central) or hypergonadotropic (i.e., secondary to gonadal failure). The underlying pathology of either type is not understood. Females usually retain their ability to become pregnant; about six successful pregnancies are described in the literature. One female had absence of the uterus [Tranebjærg, unpublished].

Symptoms to enquire about include for children, delayed puberty (the absence of secondary sexual characteristics by 14 years in a girl or 16 years in a boy), pubertal arrest. In adult men, ask about erectile impotence, reduced libido, and any history of impaired fertility or oligo/azoospermia. On examination, small, soft testes have been reported. For women, ask about amenorrhoea or oligomenorrhea, infertility loss of libido, and dyspareunia. Helpful investigations include assessment of sex hormone levels (testosterone (or oestradiol), FSH and LH, and inhibin B in males.

Management involves hormone replacement in the standard way (i.e testosteron substitution in male patients, estrogen-gestagen substitution in female patients).

**Hypothyroidism**
The frequency of thyroid dysfunction in Wolfram syndrome is not known. It is prudent to include an assessment of TSH in annual review investigations; and in the presence of symptoms, to measure free-T3, free-T4 and TSH. Thyroid substitution therapy can be given if required with L-Thyroxine (starting dose 25micrograms/day in children, 50 micrograms/day in adults).

**Growth retardation.** Most adults have normal height, but growth retardation is not infrequent. This may relate to pubertal disturbance in those with hypogonadism. Linear growth should be monitored in children using standard growth charts.
**Wolfram Syndrome guide for neurologists**

**Management of neurological involvement by neurologists or neuro-paediatricians**

In Wolfram Syndrome almost every organ/body system may be affected. Wolfram Syndrome is typically associated with sensorineural hearing loss, and other progressive neurological abnormalities. The natural history of Wolfram Syndrome was described in 45 individuals studied (mean age 16 years, range 5-32 years) from 29 families in the UK (Barrett et al 1995). Hearing impairment was present in 64%. Sixty percent of all individuals had one or more of the following signs and/or symptoms: ataxia, peripheral neuropathy, mental retardation, early onset dementia (disinhibition and/or short term memory loss), psychiatric illness (most commonly depression), and central sleep apnoea. MRI scans in individuals with this syndrome may show generalised brain atrophy with loss of the posterior pituitary bright spot, thinning of the optic nerves, and loss of volume of the cerebellum and brainstem.

**Suggested management**

Annual neurological examination for asymptomatic patients and bi-annually for symptomatic patients

Brain MRI scan at diagnosis and to be repeated if acute deterioration of neurological signs and/or symptoms or at adult age

**Cerebellar ataxia assessment:**
- Use of validated ataxia-specific rating scales for measuring progression: SARA (see supplementary data)
- Washington unified rating scale (WURS)

**Management** – Multidisciplinary team input and rehabilitation including:
- Ophthalmology services and visual impairment team input to optimise visual functioning
- Physiotherapy and occupational therapy team input with regards physical (gross motor and fine motor / coordination) difficulties
- Speech and language therapy input with regards dysarthria (speech difficulties) and swallowing difficulties (which may lead to recurrent chest infections due to aspiration)
- Drug treatments for spasticity (oral anti-spasticity medications such as baclofen, and/or botulinum toxin injections)

**Brainstem (respiratory drive) involvement assessment:**
- Screening by polysomnography or nocturnal oximetry (every 2 years)
- If symptoms: bronchoscopy (vocal cord mobility, obstructive cause), spirometry and morning blood gases

**Management** – as per respiratory / ventilation experts (tracheostomy and ventilatory support if needed)

**Peripheral neuropathy assessment:**
- Presence of symptoms such as numbness, tingling, burning, jabbing or electric-like pain or absence of deep tendon reflexes
- Presence of signs and / or symptoms of cardiovascular and / or gastrointestinal autonomic neuropathy
- Nerve conduction studies, Tilt-test in presence of autonomic cardiovascular symptoms, other investigations as per advice of cardiology and / or gastroenterology specialists

If neuropathic pain is present consider starting treatment for this – eg. Gabapentin, Pregablin, Carbamazepine, Amitriptyline, Lidocaine patch and / or transcutaneous electrical nerve stimulation (TENS)

**Epilepsy assessment:**
Electroencephalography if seizures occur
Treatment - Anti-epileptic drugs and counselling

**Cognitive assessment:**
Neuropsychological testing adapted to age (Children: WISC-IV; Adult: MMSE, FAB) and to vision difficulties

Management - Rehabilitation, special education

**Mental health assessment:**
Assessment includes taking a complete history and performing a detailed examination. Consider patient’s appearance, behaviour, speech, mood, thinking and any abnormal perceptions.

Screening for anxiety, depression, abnormal behaviour (obsessive-compulsive behaviours, aggression, eating disorders etc.) or psychosis

Management – consider referral for expert psychiatric input
Wolfram syndrome guide for ophthalmologists

The combination of insulin dependent diabetes mellitus presenting under 15 years and progressive optic atrophy is pathognomonic for Wolfram syndrome. The PPV and NPV is estimated at 85% (Barrett et al 1995).

Diagnostic criteria for optic atrophy: optic atrophy is defined as generalised pallor of the optic discs on direct fundoscopy. The presenting symptoms are decreasing visual acuity and loss of colour vision, and affected patients commonly complain of ‘everything going grey’. The optic atrophy classically occurs before 15 years of age, and is progressive, leading to reduced visual acuity less than 6/60 within a median of 8 years (ref Barrett Eye). Visual evoked responses show normal latency but very low amplitudes (Mtanda). Other ophthalmologic findings reported in WFS but not confirmed as part of the phenotype include cataracts, described in eight patients [Hansen et al 2005], and nystagmus. Optic atrophy presented in 38 patients with reduced visual acuity and colour vision defect (median age 11 years), progressing to visual acuity of 6/60 or less in 35 patients (median time 8 years, range 1-25 years). Visual field examinations recorded before acuity deteriorated showed central scotomas with peripheral constriction. Blind patients had absent pupillary reflexes. Horizontal nystagmus was seen in patients with other signs of cerebellar degeneration. There was no pigmentary retinal dystrophy; only 3 patients had background diabetic retinopathy, despite a median duration of diabetes of 24 years. Electoretinography was normal in 3 patients and showed reduced amplitude in 3 patients; visual evoked responses were abnormal (10/10 patients: reduced amplitude to both flash and pattern stimulation). Magnetic resonance imaging showed generalised brain atrophy with reduced signal from the optic nerves and chiasm. A post-mortem brain specimen from one patient revealed atrophy of the optic nerves, chiasm, cerebellum and brainstem. This primary neurodegenerative disorder presents with diabetes mellitus and progressive optic atrophy, probably due to pathology in the optic nerve.

Management:
At diagnosis: eye examination, including refraction and visual acuity, slit-lamp examination, colour vision testing, visual field (Goldman perimetry), funduscopy, OCT scan of the retinal nerve fiber layer, visual evoked potentials. Fundus auto fluorescence testing, fluorescein angiography and electroretinogram may be required in case of retinal involvement.

Correction of refractive error (myopia, hyperopia), filtrating glasses (if photoaversion)

Yearly eye examination: visual acuity, funduscopy, visual field and OCT scan are mandatory. Other tests as described at diagnosis, depending on the course of the disease

Cataract surgery if needed. Magnifying glasses, digital systems, voice systems depending of the level of visual acuity. Loss of visual field requires assistance and rehabilitation procedure for moving outside.
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Further information Wolfram Multidisciplinary Services

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Version 2 August 2014 Miss Archana Kulkarni, Mr John Ainsworth
Wolfram syndrome guide for geneticists

Wolfram syndrome (WS) (OMIM 222300) is the inherited association of childhood onset diabetes mellitus (usually before 15 years) with progressive optic atrophy (Wolfram and Wagener 1938), also known as DIDMOAD (Diabetes Insipidus, Diabetes Mellitus, Optic Atrophy and Deafness). It is a progressive neurodegenerative disorder, and many patients also develop urinary tract atony, ataxia, peripheral neuropathy, dementia and other psychiatric illnesses. Although the median age at death is 30 years, some patients have been known to survive into their 6th decade. This is an autosomal recessive monogenic disease, and most affected patients have mutations in the wfs1 gene (ref). Several families of Jordanian origin have been described with mutations in a second gene, ZCD2 (ref). These children had a subtly different phenotype without diabetes insipidus but with gastrointestinal bleeding.

Wolfram syndrome-like disorder has been proposed as a name for two families who were characterised by older onset diabetes mellitus, sensorineural hearing loss, psychiatric illness and variable optic atrophy. Both families had a missense mutation in the wfs1 gene, apparently dominantly inherited.

Finally, wfs1-related Low Frequency Sensorineural Hearing Loss (LFSHL) is also caused by mutations in the wfs1 gene. It is characterised by the autosomal dominant inheritance of congenital, non-syndromic, slowly progressive, low-frequency (<2,000Hz) sensorineural hearing loss.

WS occurs:
- In children presenting most often during the first decade of life, with a mean age of diagnosis of ~11 years. This coincides with the development of optic atrophy in addition to diabetes mellitus.
- In all races, but at a greater prevalence in those where first cousin marriages are common, reflecting the autosomal recessive inheritance of the syndrome.
- Usually without a family history of diabetes; but there may be a greater risk for psychiatric illness such as depression in first or second degree relatives (Swift 1998).
- In the presence of ketosis or ketoacidosis in a minority of patients (ref). This presentation is responsible for the misclassification of WS patients as Type 1 diabetes mellitus (T1DM).
- Without T1DM associated HLA haplotypes, and without associated islet cell autoimmunity. The underlying pathology is progressive beta cell loss probably through apoptosis.

Other conditions that may be confused with WS include:

- **Wolfram syndrome-like disorder.** There are a small number of patients who have been described with onset of diabetes mellitus and / or progressive optic atrophy in adulthood (ref Nunes, Diabetic Med paper). In the family reported by Valéro et al [2008], the 60-year-old male proband had non-insulin-dependent diabetes mellitus and congenital moderate hearing impairment (50-60 dB HL over all frequencies) that had required use of a hearing aid since childhood. His 81-year-old mother had non-insulin-dependent diabetes mellitus, mild hearing impairment (no hearing aid needed), and bilateral optic atrophy since age 60 years (i.e., WFS). No other signs of WFS were present in mother and son [Valéro et al 2008].

In the family reported by Eiberg et al [2006], autosomal dominant optic atrophy, hearing impairment, and impaired glucose regulation were observed. One individual had undiagnosed diabetes mellitus, one had impaired glucose tolerance by oral glucose tolerance test (OGTT), and others had poor pancreatic b-cell function as demonstrated by the insulinogenic index (calculated as: [the 30 minute post-OGTT serum insulin concentration minus the fasting serum insulin concentration] divided by the 30 minute post-OGTT plasma glucose concentration).

The occurrence of (milder) optic atrophy in patients/families with dominantly inherited WFS-like disorder [Eiberg et al 2006] suggests that diabetes mellitus and congenital moderate hearing
impairment in the absence of optic atrophy may be an under-recognized presentation of heterozygosity for WFS1 mutations, behaving in a dominant fashion.

- **Wolfram syndrome type 2** (WFS2) ([OMIM 604928](https://omim.org/entry/604928)), diagnosed in four Jordanian families and caused by mutations in ZCD2 on 4q22, is characterized by juvenile-onset diabetes mellitus, optic atrophy, high-frequency sensorineural hearing impairment, urinary tract dilatation, impaired renal function, hypogonadism, and severe gastrointestinal ulcer and bleeding, but not diabetes insipidus [El-Shanti et al 2000, Al-Sheyyab et al 2001, Amr et al 2007]. In one family the facial features were abnormal [Amr et al 2007]. The disorder is apparently very rare and may be confined to a certain ethnic background. Of note, molecular genetic testing of 377 hearing impaired probands did not reveal additional individuals with ZCD2 mutations, indicating that mutation of ZCD2 does not explain a substantial fraction of nonsyndromic hearing impairment.

- **WFS1-related Low Frequency Sensorineural Hearing Loss (WFS1-related LFSNHL)** Approximately 20% of genetic hearing impairment is inherited in an autosomal dominant manner, a small fraction of which is LFSNHL. Mutations in WFS1 were identified in ten families out of 13 families with autosomal dominant LFSNHL in whom linkage studies either showed linkage or were compatible with linkage to chromosome 4p [Bespalova et al 2001, Young et al 2001, Cryns et al 2002]. In five of 30 Danish families [Tranebjærg et al 2004] and three of nine Japanese families [Fukuoka et al 2007] with characteristic findings on audiogram and/or a positive family history that were unsuitable for linkage analysis, molecular genetic testing showed WFS1 mutations.

- **Mitochondrial diabetes.** Maternal transmission of mutated or deleted mitochondrial DNA can result in diabetes. The commonest (albeit rare) form is a point substitution at nucleotide position 3243 (A to G) in the mitochondrial tRNA (leu UUR) gene (van Den Ouweland 1992). This form of mitochondrial diabetes is associated with high tone sensorineural deafness and occasionally short stature. The diabetes is characterised by progressive non-autoimmune beta cell loss and insulin dependence.

- **Thiamine responsive megaloblastic anaemia syndrome** (Roger’s syndrome). This is the triad of early onset (under 5 years of age) diabetes mellitus, sensorineural deafness and megaloblastic anaemia (Rogers 1969). This is due to mutations in the SLC19A2 gene (ref Labay). The diabetes is insulin dependent, but may respond to pharmacologic doses of vitamin B1 (Thiamin). Most patients develop an insulin requirement by puberty (Ozdemir Ped Diabetes 2002).

- **Alström syndrome** is characterized by infancy onset cone-rod dystrophy and obesity. Other features include progressive sensorineural hearing impairment, dilated cardiomyopathy, severe insulin resistance, and developmental delay. Cone-rod dystrophy presents as progressive visual impairment, photophobia, and nystagmus starting between birth and age 15 months. Affected individuals have no light perception by age 20 years. Children usually have normal birth weight but become obese during their first year. Progressive sensorineural hearing loss begins in the first decade in as many as 70% of individuals. Severe insulin resistant diabetes often presents by puberty. Other endocrine abnormalities can include hypothyroidism and male hypogonadotrophic hypogonadism. Over 60% of individuals with Alström syndrome develop cardiac failure as a result of dilated cardiomyopathy at some stage of their lives. Approximately 50% of individuals have delay in early developmental milestones. Urologic disorders of varying severity, characterized by detrusor-urethral dyssynergia, appear in females in their late teens. Severe renal disease is usually a late finding. This is a monogenic, recessively inherited form of diabetes, due to mutations in the ALM1 gene.

- **Bardet-Biedl syndrome** (BBS) is characterized by cone-rod dystrophy, truncal obesity, postaxial polydactyl, cognitive impairment, male hypogonadotrophic hypogonadism, complex female
genitourinary malformations, and renal dysfunction. Birth weight is usually normal, but significant weight gain begins within the first year. Insulin resistant diabetes manifests in adolescence or adulthood. A majority of individuals have significant learning difficulties. Renal disease is a major cause of morbidity and mortality. Mutations have been found in at least twelve genes, and inheritance is autosomal recessive.

- **Friedreich ataxia** (FRDA) is characterized by slowly progressive ataxia with mean age of onset between ten and 15 years and usually before age 25 years. FRDA is typically associated with depressed tendon reflexes, dysarthria, muscle weakness, spasticity in the lower limbs, optic nerve atrophy, scoliosis, bladder dysfunction, and loss of position and vibration senses. About two thirds of individuals with FRDA have cardiomyopathy, 30% have diabetes mellitus, and approximately 25% have an "atypical" presentation with later onset, retained tendon reflexes, or unusually slow progression of disease. Mutations in FXX are causative. Inheritance is autosomal recessive.

- **Kearns-Sayre syndrome** (see Mitochondrial DNA Deletion Syndromes). Mitochondrial DNA (mtDNA) deletion syndromes comprise three overlapping phenotypes that may be observed in different members of the same family or may evolve in a given individual over time: Kearns-Sayre syndrome (KSS), Pearson syndrome, and progressive external ophthalmoplegia (PEO). Individuals with KSS have the onset of pigmentary retinopathy and PEO before age 20 years and at least one of the following: cardiac conduction block, cerebrospinal fluid protein concentration greater than 100 mg/dL, or cerebellar ataxia. Other frequent but not invariable clinical manifestations include short stature, hearing loss, dementia, limb weakness, diabetes mellitus, hypoparathyroidism, and growth hormone deficiency. Approximately 90% of individuals with KSS have a large-scale (i.e., 1.3-10 kb) mtDNA deletion that is usually present in all tissues; however, mutant mtDNA is often undetectable in blood cells, necessitating examination of muscle. When inherited, mtDNA deletion syndromes are transmitted by maternal inheritance.

- **Optic atrophy type 1** (OPA1, or Kjer type optic atrophy) (OMIM 605290, OMIM 165500) is characterized by bilateral and symmetric optic nerve pallor associated with insidious decrease in visual acuity usually between ages four and six years, visual field defects, and colour vision defects. Visual impairment is usually moderate (6/10 to 2/10), but ranges from mild or even insignificant to severe (legal blindness with acuity <1/20). Other findings can include auditory neuropathy resulting in sensorineural hearing loss that ranges from severe and congenital to subclinical (i.e., identified by specific audiologic testing only). Mutations in OPA1 are causative. Inheritance is autosomal dominant [Payne et al 2004, Amati-Bonneau et al 2005].

- **Deafness-dystonia-optic neuronopathy syndrome** (DDON, or Mohr-Tranebjærg syndrome) (OMIM 304700). Males with DDON have a progressive auditory neuropathy with prelingual or postlingual sensorineural hearing impairment, slowly progressive dystonia or ataxia in the teens, slowly progressive decreased visual acuity from optic atrophy beginning at about age 20 years, and dementia beginning at about age 40 years. Psychiatric symptoms such as personality change and paranoia may appear in childhood and progress. The neurologic, visual, and neuropsychiatric signs vary in degree of severity and rate of progression. Females may have mild hearing impairment and focal dystonia. Mutations in TIMM8A are causative. Inheritance is X-linked.

Genetic testing
Index case : WFS1 +/- CISD2 screening
1 or 2 mutated alleles : perform mutation screening in parents of index case

Genetic counseling
Information about recurrence risk to parents (25%), to adult patients and extended family members.

Prenatal Diagnosis (PN)
Available only for families in which the disease-causing mutation has been identified
For 25% recurrence risk (example: parents of an index case)

Preimplantation Genetic Diagnosis (PGD)
To discuss with referral centres (may be available for families in which the disease-causing mutation has been identified).
My name is Denise Williams and I am a doctor specialising in genetic conditions. My role in the Wolfram syndrome clinics is to help individuals and their families by providing understandable information about the condition, explaining the way in which Wolfram syndrome is inherited and arranging genetic testing if this has not already been done. If appropriate I take the opportunity to discuss the risks to future children and / or other family members. The questions I am asked are very varied – I normally suggest you write them down before you attend the clinic as it is a long day and very busy, so important things are easily forgotten. I hope to provide the information necessary to allow families to make informed medical and personal choices.

The first time I meet a family I expect to spend about three quarters of an hour with them, but some people need more and some less. Subsequent appointments are usually shorter and sometimes I am not needed. I always think that people will want to ‘dip in and out’ of genetics services at different points of their lives. What’s really important is that I offer to see the young people affected by Wolfram syndrome in their own right. Genetics is taught really well in school, most commonly in year 10 and year 11, so I like to start discussions at this time. The teenagers pick it up really easily and often ask me challenging questions!

I work closely with the genetic counsellors, Shagufta Khan in the paediatric clinics at Birmingham Children’s Hospital and Chris Platt in the adult clinics at the Queen Elizabeth Hospital in Birmingham. Both Shagufta and Chris have specialist training in genetics and counselling skills. Before your first appointment, either Shagufta or Chris will try to telephone you to obtain some basic information before you attend the clinic. They may ask about your family, so that a family tree can be drawn. They may also ask your permission to obtain relevant medical information, including the results of any genetic investigations already carried out. Having this information ahead of the clinic allows us to make the best use of the time available in clinic.

Individuals and families attending the clinic have different knowledge, depending upon their previous experiences. I nearly always go ‘back to basics’ and discuss the way in which the condition is inherited assuming no previous knowledge. I feel more comfortable doing this as I know I haven’t missed anything out and the concepts involved are quite complicated – many people find going over the same thing a second or third time is quite helpful. I use visual aids to help me; I have photographs of chromosomes and I draw diagrams illustrating the way in which genes are passed from the parents to their children. I also have some embossed charts to help me communicate effectively with individuals who have a severe visual impairment.

If I have been able to obtain the results of any genetics tests, then I will be able to discuss these in detail in clinic. If not then testing can be initiated using a small sample of blood. Genetic tests are often complex and the results can take several weeks or months. We confirm the diagnosis of Wolfram syndrome when we are able to find 2 misprints, a bit like spelling mistakes, in the ‘Wolfram syndrome’ gene. This gene is known as WFS1. In the majority of families we have been able to confirm the diagnosis, but this is certainly not true for everyone. Like many rare conditions, Wolfram syndrome is not always straightforward so it is really important that everyone is assessed individually and regular evaluation and surveillance put in place according to that individual’s needs. For example, we are starting to see some
individuals with a few of the clinical problems we expect in Wolfram syndrome, but
the pattern of these features is a bit different to those we recognise in the classic form
of the condition. Some of these individuals have a ‘misprint’ in only one copy of
their WFS1 gene and we say they have Wolfram syndrome-like disease. This seems
to be very uncommon, but is important because it is inherited in a different way to the
classic form of the condition. When this is important for a family I always discuss
this carefully.

As mentioned already, part of my role in clinic is to discuss the ‘risks’ to other family
members. From a practical point of view, finding the specific misprints responsible
for causing the Wolfram syndrome in an individual means that we can use this
information to test other family members who may be at risk of the condition or wish
to know if they are carriers. I normally suggest this is arranged with the advice and
support of the local clinical genetics service and I am in a position to liaise with my
colleagues around the country and facilitate this.

Although there is currently no cure for Wolfram syndrome and this may still be a long
way off, Professor Barrett who leads the multidisciplinary team in Birmingham is
working closely with research doctors across Europe and America. There is no doubt
that a greater understanding about the different misprints in the gene and the different
ways in which people are affected by the condition, may lead to novel treatments for
different aspects of it in the future. This is something we are all hoping to discuss
with you in time.
Wolfram syndrome guide for urologists

Management of urological involvement by urologists, rehabilitation physicians and neurologists

Urge incontinence due to bladder syssynergia or neuropathic bladder has been reported in up to about 60% of affected people. These features can present during childhood. The pathology is not well understood, but it is thought that there is a loss of upper motor neurone control over the detrusor muscle; in addition, autonomic nervous system dysfunction may lead to altered bladder sensation. Reports from case series show pelviureteric dilatation on ultrasound scans. The part played by cranial diabetes insipidus is unclear.

Baseline investigations
- Ask about urinary symptoms, complete a voiding diary, undertake a clinical examination for evidence of neurological involvement.
- Check renal function (blood electrolytes, urea, creatinine, glomerular filtration rate (GFR))
- Bladder and renal ultrasound (residual urine)
- Urodynamic testing to include flow rates on emptying the bladder, and residual bladder volume.

Management may involve:
Treatment options include advice about double voiding technique; electrical stimulation; anticholinergic drugs to stabilise the detrusor muscle; botulinum toxin; clean intermittent self-catheterization; permanent indwelling catheter; ileal conduit surgery.

Screening urinary infections
Urine culture if fever or other symptoms

Intermittent self-catheterization
Preliminary assessment of the ability to self-catheterization, taking into account ataxia, low vision or cognitive deficiency (PP-Test)

Indwelling urinary catheter
Risk factors for bladder tumors

Sexual dysfunction
Management in standard way
Wolfram syndrome guide for paediatricians and other health care professionals

Version 2.

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Wolfram syndrome (WS) (OMIM 222300) is the inherited association of childhood onset diabetes mellitus (usually before 15 years) with progressive optic atrophy (Wolfram and Wagener 1938), also known as DIDMOAD (Diabetes Insipidus, Diabetes Mellitus, Optic Atrophy and Deafness). It is a progressive neurodegenerative disorder, and many patients also develop urinary tract atony, ataxia, peripheral neuropathy, dementia and other psychiatric illnesses. Although associated with a shorter life span, some patients have been known to survive into their 6th decade. This is an autosomal recessive monogenic disease, and most affected patients have mutations in the WFS1 gene.

Wolfram syndrome-like disorder has been proposed as a name for two families who were characterised by older onset diabetes mellitus, sensorineural hearing loss, psychiatric illness and variable optic atrophy. Both families had a missense mutation in the WFS1 gene, apparently dominantly inherited.

Finally, WFS1-related Low Frequency Sensorineural Hearing Loss (LFSNHL) is also caused by mutations in the WFFS1 gene. It is characterised by the autosomal dominant inheritance of congenital, non-syndromic, slowly progressive, low-frequency (<2,000Hz) sensorineural hearing loss.

WS occurs:

- In children presenting most often during the first decade of life, with a mean age of diagnosis of ~11 years. This coincides with the development of optic atrophy in addition to diabetes mellitus.
- In all races, but at a greater prevalence in those where first cousin marriages are common, reflecting the autosomal recessive inheritance of the syndrome.
- Usually without a family history of diabetes; but there may be a greater risk for psychiatric illness such as depression in first or second degree relatives.
- In the presence of ketosis or ketoacidosis in a minority of patients. This presentation is responsible for the misclassification of WS patients as Type 1 diabetes mellitus (T1DM).
- Without T1DM associated HLA-haplotypes, and without associated islet cell autoimmunity. The underlying pathology is progressive beta cell loss probably through apoptosis.

Other conditions that may be confused with WS include:

- **WFS1-related Low Frequency SensoriNeural Hearing Loss (WFS1-related LFSNHL)** Approximately 20% of genetic hearing impairment is inherited in an autosomal dominant manner, a small fraction of which is LFSNHL. Mutations in WFS1 were identified in ten out of 13 families with autosomal dominant LFSNHL in whom linkage studies either showed linkage or were compatible with linkage to chromosome 4p. In five of 30 Danish families and three of nine Japanese families with characteristic findings on audiogram and/or a positive family history that were unsuitable for linkage analysis, molecular genetic testing showed heterozygous WFS1 mutations.

- **Wolfram syndrome-like disorder.** There are a small number of patients who have been described with onset of diabetes mellitus and / or progressive optic atrophy in adulthood. In the family...
reported by Eiberg et al [2006], autosomal dominant optic atrophy, hearing impairment, and impaired glucose regulation were observed. The occurrence of (milder) optic atrophy in patients/families with dominantly inherited WFS-like disorder suggests that diabetes mellitus and congenital moderate hearing impairment in the absence of optic atrophy may be an under-recognized presentation of heterozygosity for WFS1 mutations, behaving in a dominant fashion.

- **Wolfram syndrome type 2** (WFS2) (OMIM 604928), diagnosed in four Jordanian families and caused by mutations in CISD2 on chromosome 4q22, is characterized by juvenile-onset diabetes mellitus, optic atrophy, high-frequency sensorineural hearing impairment, urinary tract dilatation, impaired renal function, hypogonadism, and severe gastrointestinal ulcer and bleeding, but not diabetes insipidus. The disorder is apparently very rare and may be confined to a certain ethnic background. Of note, molecular genetic testing of 377 hearing impaired people did not reveal additional individuals with CISD2 mutations, indicating mutations in this gene do not explain a substantial fraction of nonsyndromic hearing impairment.

- **Mitochondrial diabetes.** Maternal transmission of mutated or deleted mitochondrial DNA can result in diabetes. The commonest (albeit rare) form is a point substitution at nucleotide position 3243 (A to G) in the mitochondrial tRNA (leu UUR) gene. This form of mitochondrial diabetes is associated with high tone sensorineural deafness and occasionally short stature. The diabetes is characterised by progressive non-autoimmune beta cell loss and insulin dependence.

- **Thiamine responsive megaloblastic anaemia syndrome** (Roger’s syndrome). This is the triad of early onset (under 5 years of age) diabetes mellitus, sensorineural deafness and megaloblastic anaemia (Rogers 1969). This is due to mutations in the SLC19A2 gene. The diabetes is insulin dependent, but may respond to pharmacologic doses of vitamin B1 (Thiamin). Most patients develop an insulin requirement by puberty.

- **Alström syndrome** is characterized by infancy onset cone-rod dystrophy and obesity. Other features include progressive sensorineural hearing impairment, dilated cardiomyopathy, severe insulin resistance, and developmental delay. Cone-rod dystrophy presents as progressive visual impairment, photophobia, and nystagmus starting between birth and age 15 months. Affected individuals have no light perception by age 20 years. Children usually have normal birth weight but become obese during their first year. Progressive sensorineural hearing loss begins in the first decade in as many as 70% of individuals. Severe insulin resistant diabetes often presents by puberty. Other endocrine abnormalities can include hypothyroidism and male hypogonadotrophic hypogonadism. Over 60% of individuals with Alström syndrome develop cardiac failure as a result of dilated cardiomyopathy at some stage of their lives. Approximately 50% of individuals have delay in early developmental milestones. Urologic disorders of varying severity, characterized by detrusor-urethral dyssynergia, appear in females in their late teens. Severe renal disease is usually a late finding. This is a monogenic, recessively inherited form of diabetes, due to mutations in the **ALMS1** gene.

- **Bardet-Biedl syndrome** (BBS) is characterized by rod-cone dystrophy, truncal obesity, postaxial polydactyly, cognitive impairment, male hypogonadotrophic hypogonadism, complex female genitourinary malformations, and renal dysfunction. Birth weight is usually normal, but significant weight gain begins within the first year. Insulin resistant diabetes manifests in adolescence or adulthood. A majority of individuals have significant learning difficulties. Renal disease is a major cause of morbidity and mortality. Mutations have been found in at least nineteen genes, and inheritance is autosomal recessive.

- **Friedreich ataxia** (FRDA) is characterized by slowly progressive ataxia with mean age of onset between ten and 15 years and usually before age 25 years. FRDA is typically associated with depressed tendon reflexes, dysarthria, muscle weakness, spasticity in the lower limbs, optic nerve atrophy, scoliosis, bladder dysfunction, and loss of position and vibration senses. About two thirds
of individuals with FRDA have cardiomyopathy, 30% have diabetes mellitus, and approximately 25% have an “atypical” presentation with later onset, retained tendon reflexes, or unusually slow progression of disease. Mutations in FXN are causative. Inheritance is autosomal recessive.

- **Kearns-Sayre syndrome** (see Mitochondrial DNA Deletion Syndromes). Mitochondrial DNA (mtDNA) deletion syndromes comprise three overlapping phenotypes that may be observed in different members of the same family or may evolve in a given individual over time: Kearns-Sayre syndrome (KSS), Pearson syndrome, and progressive external ophthalmoplegia (PEO). Individuals with KSS have the onset of pigmentary retinopathy and PEO before age 20 years and at least one of the following: cardiac conduction block, cerebrospinal fluid protein concentration greater than 100 mg/dL, or cerebellar ataxia. Other frequent but not invariable clinical manifestations include short stature, hearing loss, dementia, limb weakness, diabetes mellitus, hypoparathyroidism, and growth hormone deficiency. Approximately 90% of individuals with KSS have a large-scale (i.e., 1.3-10 kb) mtDNA deletion that is usually present in all tissues; however, mutant mtDNA is often undetectable in blood cells, necessitating examination of muscle. When inherited, mtDNA deletion syndromes are transmitted by maternal inheritance.

- **Optic Atrophy type 1** (OPA1, or Kjer type optic atrophy) (OMIM 605290, OMIM 165500) is characterized by bilateral and symmetric optic nerve pallor associated with insidious decrease in visual acuity usually between ages four and six years, visual field defects, and color vision defects. Visual impairment is usually moderate (6/10 to 2/10), but ranges from mild or even insignificant to severe (legal blindness with acuity <1/20). Other findings can include auditory neuropathy resulting in sensorineural hearing loss that ranges from severe and congenital to subclinical (i.e., identified by specific audiologic testing only). Mutations in OPA1 are causative. Inheritance is autosomal dominant.

- **Deafness-dystonia-optic neuronopathy syndrome** (DDON, or Mohr-Tranebjærg syndrome) (OMIM 304700). Males with DDON have a progressive auditory neuropathy with prelingual or postlingual sensorineural hearing impairment, slowly progressive dystonia or ataxia in the teens, slowly progressive decreased visual acuity from optic atrophy beginning at about age 20 years, and dementia beginning at about age 40 years. Psychiatric symptoms such as personality change and paranoia may appear in childhood and progress. The neurologic, visual, and neuropsychiatric signs vary in degree of severity and rate of progression. Females may have mild hearing impairment and focal dystonia. Mutations in TIMM8A are causative. Inheritance is X-linked.

**Genetic testing**
- Index case: WFS1 +/- CISD2 screening
- 1 or 2 mutated alleles: perform mutation screening in parents of index case

**Genetic counselling**
- Information about recurrence risk to parents (25%), to adult patients and extended family members.

**Prenatal Diagnosis (PN)**
- Available only for families in which the disease-causing mutation has been identified
- For 25% recurrence risk (example: parents of an index case)

**Preimplantation Genetic Diagnosis (PGD)**
- To discuss with referral centres (may be available for families in which the disease-causing mutation has been identified).
Washington University School of Medicine has a long history of pioneering medical research, including the discovery of the Wolfram syndrome gene (WFS1) and its function, led by the late Alan Permutt, MD, and his team of researchers. Today, a collaborative effort by Washington University School of Medicine faculty, led by Fumihiko (Fumi) Urano, MD, the Samuel E. Schechter Professor of Medicine, is advancing the understanding of the progression of Wolfram syndrome with the goal of identifying targets for therapeutic interventions and treatments for Wolfram syndrome.

AT WASHINGTON UNIVERSITY MEDICAL CENTER IN ST. LOUIS, USA

- Identified an enzyme implicated in endoplasmic reticulum stress as a molecular target for Wolfram syndrome treatment.
- Uncovered FDA-Approved drugs that block activation this enzyme and cell death by induced pluripotent stem cells (iPSCs) derived from patient skin cells. These pluripotent stem cells will help in the identification of therapeutics to treat Wolfram syndrome and may eventually be used to replace damaged tissues, including pancreatic β cells, brain cells, and eye cells, resulting from this monogenic disorder.
- Identified potential biomarkers reflecting the progression of the disease using blood samples from patients and their siblings.
- Several biotech and pharmaceutical companies have been identified with overlapping interests in advancing drugs of potential benefit to patients with Wolfram syndrome. With the aid of the Jack and JT Snow Scientific Research Foundation, Dr. Urano and Dr. Timothy Barrett are planning joint clinical trials.

These translational discoveries utilized an important pediatric patient population, combining multiple assessment methods and resources drawn from a dedicated basic science and clinical community working collaboratively to understand the molecular mechanism and identify actionable targets for the treatment of Wolfram Syndrome.

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http://www.erstress.com/
http://wolframsyndrome.blogspot.com/
June 2016 saw co-founders Tracy & Paul receive the Prime Minister’s Points of Light Award.

Prime Minister David Cameron said:

Since Jennifer’s diagnosis, Paul and Tracy have taken phenomenal steps to raise awareness and find a cure for Wolfram Syndrome. Their work is having an impact in the medical community and has the potential to transform the lives of those living with this rare condition. I’m pleased to name Paul and Tracy the 555th and 556th UK Points of Light.”

Tracy and Paul’s local MP, Sir Peter Bottomley MP said:

“Paul and Tracy have built a charity in Worthing that is supporting people across the country living with this rare condition. I’m delighted that their achievements are being recognised by the Prime Minister with this Point of Light award.”

https://www.pointsoflight.gov.uk/working-wolfram-syndrome/

WSUK are always pleased to hear from families and those affected by the condition.

Donations and contributions of support are always gratefully received so that we may continue to provide information, support and help fund research into trying to find medication to provide the best treatment.

Donations can be made via post or online by going to our website and following one of the links there.
About Wolfram Syndrome………

The first signs of someone being affected by WS are juvenile onset Diabetes Mellitus and Optic Atrophy (reduced vision). Some patients go on to develop hearing loss and Diabetes Insipidus (water diabetes).

These four conditions are the main features of WS, also known as DIDMOAD. There are also other health problems for those with WS which can include irregular breathing, loss of the sense of smell, depression, loss of the gag reflex and impulsive and aggressive behaviour to name but a few. Not everyone is affected the same or develops all the features. WS affects 1 in 770,000 people in the UK.

Research

Research is being carried out at Birmingham Children’s Hospital and The Queen Elizabeth Hospital in Birmingham, as well as at other hospitals and universities around the world. We help to provide funding towards the research carried out here in the UK and will shortly be helping to fund the drug trial due to start in the Autumn of 2018.

WSUK keeps up to date with the medical research and is in contact with medical experts who can offer advice.

We maintain the only UK database of those affected by this disorder.

We are endeavouring to get information to hospitals and GP practices around the UK.

How it all started

Wolfram Syndrome is a rare, progressive neurodegenerative condition, which is life shortening.

Wolfram Syndrome UK (WSUK) is the only national charity and support group to help fund research and provide support for those affected by the condition and their families in the UK.

The support group and website were started in 2010 by Paul & Tracy Lynch from Worthing, West Sussex after their daughter, then aged eight, was diagnosed with WS. The only website associated with WS then was a worldwide site. There was no easy to read information available and no real support, as many medical professionals had or still have never heard or come across the syndrome.

WSUK became a registered charity in June 2013. Prior to that we had been raising funds for research via WellChild, the national charity for sick children, and our fundraising group ‘The Charity Roadtrip’. WSUK and WellChild continue to work closely together.

What we are doing

There is a dedicated UK telephone line and website giving up to date information on events, research, trials, along with links to other useful sites, multi disciplinary clinics and Social Media pages for anyone linked to WS to post on. We hold an annual conference for families & doctors to receive useful information and to just meet up. We also send out a quarterly newsletter with news of fundraising events and other useful information.

WSUK are working in alliance with WellChild and Birmingham Children’s Hospital to raise awareness, bring support to patients and families and to advance research into trying to find a way of halting or slowing down the progression of this syndrome.

Research is being carried out at Birmingham Children’s Hospital and The Queen Elizabeth Hospital in Birmingham, as well as at other hospitals and universities around the world. We help to provide funding towards the research carried out here in the UK and will shortly be helping to fund the drug trial due to start in the Autumn of 2018.

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Management of Wolfram Syndrome

A Clinical Guideline

Wolfram Syndrome Guideline Development Group
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Introduction...

... to Wolfram Syndrome
Wolfram syndrome (WS), also known as DIDMOAD (Diabetes Insipidus, Diabetes Mellitus (DM), Optic Atrophy (OA), and Deafness) is a rare autosomal recessive disorder. The estimated prevalence of WS is 1 in 770,000. The minimal criteria for diagnosis are juvenile-onset DM and OA but patients may also develop diabetes insipidus, sensorineural deafness, renal tract abnormalities, and neuropsychiatric disorders; and variants exist with only partial features. The prognosis is mainly linked to the severity of the neurological symptoms.

WS is a genetically heterogeneous disease. Most patients carry mutations in the WFS1 gene, encoding an endoplasmic reticulum membrane embedded protein called Wolframin. CISD2 is a second causative gene associated with WS. It encodes a mitochondrial protein.

In addition, mutations in the WFS1 gene are also associated with the poorly defined ‘Wolfram-Like Syndrome (WS-like) disorders’ including DM, OA, or deafness in dominant or recessive families, and in dominantly-inherited low-frequency sensorineural hearing loss (LFSNHL).

... to the Wolfram syndrome guideline project
These guidelines have been developed by referring physicians involved in the EURO-WABB project, according to the DYSCERNE guideline development process (www.dyscerne.org.dysc.home/). The experts who participated in the guideline development are listed on page 17.

... to the Wolfram syndrome clinical management guidelines
What are the aims of the guidelines?
The guidelines aim to provide recommendations for the diagnosis, management and the follow-up of patients with WS. As it is a multisystemic disorder, WS patients may require various tests, screening and multidisciplinary interventions at different stages of their lives. These recommendations aim to support high quality care for people with WS in a format that is accessible to anybody who is involved in the care of these patients. Note that transition is a process which includes the event of transfer from childrens’ to adult services and needs to attend to the medical, psychosocial, and educational/vocational needs of the young person and his/her parents/carers. Care needs to be provided that includes attention to transition needs.

How are they organised?
The guidelines are divided into
- clinical features and diagnostic criteria
- baseline investigations
- recommended tests, that are listed and organised into specific groups corresponding to the different symptoms and affected organs. Any recommendations that are specifically addressed either to children or to adult patients are specified.

A list of references starts on page 14, organised according to the different sections of the guidelines. Additionally, there is a list of useful contacts for patients and families affected by WS, on page 18.

Note: N=normal; ABNL=abnormal
## Diagnosis and clinical features of Wolfram Syndrome

### Diagnostic criteria of WS

<table>
<thead>
<tr>
<th>Major criteria</th>
<th>Minor criteria</th>
<th>Minimum required</th>
<th>Other variable suggestive evidence:</th>
</tr>
</thead>
<tbody>
<tr>
<td>- Diabetes mellitus &lt;16 yrs &lt;br&gt; - Optic atrophy &lt;16 yrs</td>
<td>- Diabetes insipidus &lt;br&gt; - Diabetes mellitus &gt; 16 yrs &lt;br&gt; - Optic atrophy &gt;16 yrs &lt;br&gt; - Sensorineural deafness &lt;br&gt; - Neurological signs (ataxia, epilepsy, neuropathy, cognitive impairment) &lt;br&gt; - Renal tract abnormalities &lt;br&gt; - 1 loss of function mutation in WFS1/CISD2 AND/OR family history of Wolfram syndrome</td>
<td>-2 major OR -1 major plus 2 minor criteria</td>
<td>- Hypogonadism (males) &lt;br&gt; - Absence of type 1 diabetes auto-antibodies &lt;br&gt; - Bilateral cataracts &lt;br&gt; - Psychiatric disorder &lt;br&gt; - Gastrointestinal disorders</td>
</tr>
</tbody>
</table>

Table 1: Diagnostic criteria. Note: The diagnosis is established in individuals of all ages in whom two pathological WFS1 or CISD2 mutations are identified.

### Wolfram Syndrome-like disorders: variable mode of inheritance

At least 1 criterion among diabetes mellitus (or glucose intolerance), optic atrophy or deafness

AND

At least one loss of function WFS1 or CISD2 mutation

### Differential diagnosis includes:

- Mitochondrial disorders: Maternally Inherited Diabetes mellitus and Deafness, Leber Hereditary Optic Neuropathy
- Thiamine-responsive megaloblastic anemia, diabetes and deafness
- Autosomal Dominant Optic Atrophy
- X-linked Charcot-Marie-Tooth disease type 5
- Deafness, Dystonia, Optic Neuropathy syndrome
- Friedreich ataxia
- Bardet-Biedl syndrome
- Alstrom syndrome
# Recommended baseline investigations in Wolfram Syndrome

<table>
<thead>
<tr>
<th>Clinical Features of WS</th>
<th>Baseline investigations</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Endocrine system</strong></td>
<td>Fasting plasma glucose and HbA1c. Type 1 diabetes associated auto-antibodies most often absent: mainly glutamate decarboxylase (GAD), tyrosin phosphatase (IA-2) and insulin antibodies, if available islet cell Ab (ICA) or ZnT8 Ab. Low insulin reserve assessed by basal and/or post standard meal stimulated C- Peptide measurements. <em>Note that Wolfram patients present rarely with diabetic ketoacidosis.</em></td>
</tr>
<tr>
<td>… Diabetes Mellitus</td>
<td>Morning paired urine and fasting plasma for osmolarity and sodium concentration after nocturnal and morning euglycaemia.</td>
</tr>
<tr>
<td>… Diabetes Insipidus</td>
<td>Testosterone, FSH and LH, inhibin B</td>
</tr>
<tr>
<td>… Hypogonadism (male)</td>
<td>Visual acuity, fundus examination, visual field, OCT scan, visual evoked potentials, colour vision testing</td>
</tr>
<tr>
<td><strong>Sensory involvement</strong></td>
<td>Audiogram, auditory evoked potentials</td>
</tr>
<tr>
<td>… Optic Atrophy</td>
<td>Neurological examination with brain MRI and cognitive assessment. Other specific investigations according to the results of clinical examination. Mental health assessment. Consider test of olfaction</td>
</tr>
<tr>
<td>… Hearing Loss</td>
<td>Questionnaire regarding urinary symptoms with voiding diary, Assessment of renal function (blood electrolytes, urea, creatinine, GFR), ultrasound renal tract and urodynamic testing.</td>
</tr>
<tr>
<td><strong>Neurological signs</strong></td>
<td><strong>Confirmation of WS diagnosis</strong></td>
</tr>
<tr>
<td><strong>Urological signs</strong></td>
<td><strong>Molecular Analysis</strong></td>
</tr>
<tr>
<td>WFS1. Analysis of CISD2 only if negative WFS1 screening, characteristic phenotype, or middle eastern origin</td>
<td></td>
</tr>
</tbody>
</table>
### Recommendations for the management of Wolfram Syndrome

**Endocrine System – Diabetes Mellitus (I)**

#### Diagnostic criteria of diabetes

<table>
<thead>
<tr>
<th>Criteria</th>
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<tbody>
<tr>
<td>Fasting (at least 8 hours) Plasma Glucose (FPG) ≥ 7.0 mmol/L</td>
</tr>
<tr>
<td>Casually measured Plasma Glucose (PG) ≥ 11.1 mmol/L + symptoms of diabetes (polyuria, polydipsia and unexplained weight loss)</td>
</tr>
<tr>
<td>2 hour PG ≥ 11.1 mmol/L in a 75-g oral glucose tolerance test</td>
</tr>
</tbody>
</table>

*If there are no osmotic symptoms or ketone production, then a confirmatory glucose test must be done on another day. In a child, raised glucose measurement should lead to same day referral to a hospital specialist experienced in management of childhood diabetes and should not delay initiation of treatment to avoid rapid deterioration (diabetic ketoacidosis: DKA)*

#### Management of DM for children by an interdisciplinary pediatric diabetes healthcare team

<table>
<thead>
<tr>
<th>Category</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Intensive education</strong></td>
<td>Insulin injection, dosage adjustment, blood glucose and ketone testing, exercise, nutrition, formal smoking avoidance, prevention and management of DKA and hypoglycemia.</td>
</tr>
<tr>
<td><strong>Glycemic targets</strong></td>
<td>Improve metabolic control to reduce diabetes-related complications with strategies tailored to each child, according to individual risk factors and vulnerability to severe hypoglycemia. HbA1c goals should be &lt;7.5%. Less stringent A1C goals may be appropriate for patients with a history of severe hypoglycemia, limited life expectancy, advanced microvascular or macrovascular complications, or extensive comorbid conditions.</td>
</tr>
<tr>
<td><strong>Insulin therapy</strong></td>
<td>Insulin regimen chosen according on age, duration of diabetes, lifestyle, socioeconomic factors, and family, patient and physician preferences. Intensive management is usually required: continuous subcutaneous insulin infusion or multiple daily injection regimens using basal insulin analogues.</td>
</tr>
<tr>
<td><strong>Glucose monitoring</strong></td>
<td>Self-monitoring of blood glucose (adapted devices for vision impaired people), glucose diary, and quarterly HbA1c measurement. If necessary and available, Continuous Glucose Monitoring System (CGMS) can be used.</td>
</tr>
<tr>
<td><strong>Nutrition</strong></td>
<td>Regular evaluation (at least annually) with nutrition counseling (based on the nutritional needs, eating habits, lifestyle, ability and interest) ensuring normal growth and development with optimal glycaemic control.</td>
</tr>
<tr>
<td><strong>Hypoglycemia</strong></td>
<td>Significant risk of hypoglycemia often necessitates less stringent glycemic goals or the use of a continuous glucose monitoring system. Severe hypoglycemia should be treated with intravenous dextrose (hospital) or subcutaneous glucagon (at home) followed by buccal glucose syrup.</td>
</tr>
</tbody>
</table>
Recommendations for the management of Wolfram Syndrome

*Endocrine System – Diabetes Mellitus (II)*

<table>
<thead>
<tr>
<th>Management of DM for children by an interdisciplinary pediatric diabetes healthcare team</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Chronic poor metabolic control</strong></td>
</tr>
<tr>
<td><strong>DKA</strong></td>
</tr>
<tr>
<td><strong>Psychological issues</strong></td>
</tr>
</tbody>
</table>
### Recommendations for the management of Wolfram Syndrome

**Endocrine System – Diabetes Mellitus (III)**

<table>
<thead>
<tr>
<th>Management of diabetes complications</th>
<th>Details</th>
</tr>
</thead>
</table>
| **Nephropathy**                      | - Yearly screening, starting at 12 years of age, in patients with duration of diabetes >5 years  
- First morning or random urine albumin to creatinine ratio, and microalbuminuria demonstrated.  
- Introduce renoprotection with angiotensin converting enzyme (ACE) inhibitors or angiotensin receptor blockers (ARBs) as soon as microalbuminuria is confirmed. |
| **Retinopathy**                      | - Yearly screening in patients with duration of diabetes more than 5 years  
- Fundoscopy, OCT scan and fluorescein angiography if signs of diabetic retinopathy are present |
| **Neuropathy**                       | - Yearly neurological exam to look for numbness, pain, cramps and paresthesia (cf. neurological section)  
- Nerve conduction studies and dysautonomia assessment in presence of clinical signs or symptoms  
- Treat symptoms |
| **Dyslipidemia**                     | - Screen at 12 and 17y (when stabilized), or <12y if risk factors exist (obesity, familial hypercholesterolaemia)  
- Fasting total cholesterol, high-density and low-density lipoprotein cholesterol, triglycerides  
- Lipid lowering drug therapy |
| **Hypertension**                     | - Screen at least annually, use appropriate cuff size, +/- 24 hour ambulatory blood pressure monitoring  
- Lifestyle modification and anti-hypertensive drug therapy |
Recommndations for the management of Wolfram Syndrome

**Endocrine System – Others**

### Diabetes insipidus
- Symptoms to seek: polyuria and polydipsia (could be masked by the polyuria induced by poor glycemic control)
- Assessment of concentrating ability of the urine: morning paired urine and fasting plasma for osmolarity and sodium concentration – even if the patient denies symptoms. Prerequisite for the evaluation of morning urine osmolarity: nocturnal and morning euglycaemia (blood glucose levels beneath the renal threshold)
- Follow up and management in standard way (according to criteria for desmopressin administration)

### Hypo or hypergonadotropic hypogonadism
- Symptoms to seek:
  - Boys and girls: delayed puberty or pubertal arrest
  - Male adolescents and men: impaired fertility, oligo/azoospermia, erectile dysfunction, reduced libido, testicular hypotrophy
  - Women: a/oligomenorrhea, infertility, loss of libido, dyspareunia,
- Hormone levels: testosterone (or oestradiol), FSH and LH, inhibin B
- Management in standard way (*i.e.* testosterone replacement in male patients with testosterone enanthate gradually increasing 50-250mg i.m. every 3-4 weeks at age less than 18 years; alternatively testosterone undecanoate i.m.every 3 months or testosterone gel 50mg/day at age over 18 years. Oestrogen-gestagen replacement in female patients)

### Hypothyroidism
- Free-T3, free-T4 and TSH if presence of symptoms
- Thyroid substitution therapy with L-Thyroxine (starting dose 25μg/day)

### Growth retardation
- Monitoring of linear growth in children using standard growth charts
Recommendations for the management of Wolfram Syndrome

**Sensory involvement**

### Visual assessment

**At diagnosis**
- Eye examination, including refraction and visual acuity, slit-lamp examination, color vision testing, visual field (Goldman perimetry), funduscopy, OCT scan of the retinal nerve fiber layer, visual evoked potentials, systematic retinography. Fundoscopy and OCT scan if signs of diabetic retinopathy are present. Flourescin angiography could be discussed according to the severity of retinal involvement.
- Correction of refractive error (myopia, hyperopia, astigmatism).

**Follow up**
- Yearly eye examination: visual acuity, funduscopy, visual field and OCT scan are mandatory. Other tests as described at diagnosis, depending on the course of the disease.
- Cataract surgery if needed. Magnifying glasses, digital systems, voice systems depending on the level of visual acuity. Loss of visual acuity requires support from vision impairment specialists.

### Hearing assessment

**At diagnosis**
- Audiogram
- Auditory evoked potentials

**Follow up**
- Test every 2 years
- Hearing Loss
- Management with hearing aids
# Recommendations for the management of Wolfram Syndrome

**Neuro-psychiatric involvement**

**Management of neurological involvement by adult or paediatric neurologists**

<table>
<thead>
<tr>
<th>Neurologic examination yearly for asymptomatic patients and twice a year for symptomatic patients</th>
<th>Brain MRI to repeat if acute aggravation of central disorders or at adult age</th>
</tr>
</thead>
</table>
| **Cerebellar ataxia assessment**  
- Use of validated ataxia-specific rating scales for measuring progression (E.g. SARA: [http://www.neurology.org/content/suppl/2006/06/07/66.11.1717.DC1/E1.doc](http://www.neurology.org/content/suppl/2006/06/07/66.11.1717.DC1/E1.doc))  
- Therapy or rehabilitation for:  
  - Nystagmus (if disability),  
  - Cerebellar intention tremor (drug, physiotherapist, intervention),  
  - Dysarthria and swallowing disorder (swallowing therapy by speech therapist), prevention of pulmonary aspiration disease (pulmonary infection)  
| **Brainstem involvement assessment**  
- Screening by polysomnography or nocturnal oximetry (every 2 years)  
- If symptoms: bronchoscopy (vocal cord mobility, obstructive cause), spirometry, morning blood gases  
- Management in standard way by respiratory physician (tracheostomy, optimal ventilation)  
| **Central respiratory failure**|
| **Peripheral neuropathy assessment**  
- Symptoms to seek (numbness, tingling, burning, jabbing or electric-like pain) or arreflexia  
- Consider cardiovascular and gastrointestinal autonomic neuropathy  
| **Epilepsia assessment**  
- Electromyography, tilt-test in presence of symptoms, then if acute aggravation:  
- Treatment for relieve the pain (Anti-epileptic, antidepressants, lidocaine patch, TENS) or hypotension  
- Electroencephalography (EEG) if seizures occur  
- Anti-epileptic drugs  
| **Cognitive assessment**  
- Neuropsychological testing adapted to age (Children: WISC-IV) and to low vision  
- Review yearly if cognitively impaired. Rehabilitation, special education  
| **Mental health assessment**  
- Screening: anxiety, depression, abnormal behavior (compulsive aggression, eating disorders) or psychosis  
- Examine: complete history, appearance, behaviour, speech, mood, thinking, abnormal perceptions  
- Management in standard way by psychiatric expert  

SARA: Scale for the assessment and rating of ataxia;  
WISC-IV: Wechsler Intelligence Scale for Children, Fourth Edition; MMSE:Mini Mental State Examination; FAB: Frontal Assessment Battery
Recommendations for the management of Wolfram Syndrome

**Urological involvement**

Management of urological involvement by urologists, rehabilitation physicians and neurologists

<table>
<thead>
<tr>
<th>Baseline investigations</th>
<th>Standardised questionnaire regarding urinary symptoms and voiding diary, clinical examination</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Assessment of renal function (blood electrolytes, urea, creatinine, glomerular filtration rate (GFR))</td>
</tr>
<tr>
<td></td>
<td>Bladder and renal ultrasound (residual urine), urodynamic testing</td>
</tr>
</tbody>
</table>

Yearly assessment:
- Questionnaire regarding urinary symptoms and voiding diary
- Assessment of renal function (urea, creatinine, GFR)
- Bladder and renal ultrasound (PVR)

Urodynamic testing: yearly
- Clinical exam, questionnaire regarding urinary symptoms and quality of life scale twice a year
- Management in standard way according expert’s decision:
  +/− Intra-venous urography, retrograde urethrocystography (voiding), renal scintigraphy
  +/− treatment (anticholinergic drugs, botulinum toxin, intermittent self-catheterization) …
- Electrical stimulation and physiotherapy, surgical intervention when needed

<table>
<thead>
<tr>
<th>Screening urinary infections</th>
<th>Urine culture if fever or other symptoms</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intermittent self-catheterization</td>
<td>Preliminary assessment of the ability to self-catheterize, taking into account ataxia, low vision or cognitive deficiency</td>
</tr>
<tr>
<td>Indwelling urinary catheter</td>
<td>Risk factors for infection</td>
</tr>
</tbody>
</table>

Risk factors for infection
Genetic testing

- Index case: *WFS1* +/- *CISD2* screening if desired by patient or parents.
- 1 or 2 mutated alleles: perform mutation screening in parents of index case.

Genetic counselling

- Information about recurrence risk to parents (25%), to adult patients and extended family members.

Prenatal Diagnosis (PN)

- Available only for families in which the disease-causing mutation has been identified.
- For 25% recurrence risk (example: parents of an index case).

Preimplantation Genetic Diagnosis (PGD)

- To discuss with referral centres (may be available for families in which the disease-causing mutation has been identified).
1. INTRODUCTION

2. DIABETES

3. NEUROLOGICAL SIGNS

4. GENETICS
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Information for patients

Sources of information and support

The groups listed below are useful sources of support and information

- **Association du syndrome de Wolfram** ([http://asso.orpha.net/ASW/](http://asso.orpha.net/ASW/))
  Contact: Tél. +33.2.97.61.42.37  Email: nolwenn.jaffre@voila.fr

- **EURO-WABB project** – [www.euro-wabb.org](http://www.euro-wabb.org)
  The general objective of this project is to support efficient diagnosis, treatment, and research for Wolfram, Alström, Bardet-Biedl (WABB) and other rare syndromes. The project is managed by a collaboration of scientists, clinicians, and patient groups. The website contains useful information about these rare diseases, some of it in several European languages.

- **Orphanet** ([www.orpha.net](http://www.orpha.net))
  Orphanet is an online database of rare diseases and related services provided through Europe. It contains information on over 5,000 conditions and lists specialised clinics, diagnostic tests, patient and organizations, research projects and clinical trials

- **OMIM** ([http://www.omim.org/](http://www.omim.org/))
  OMIM is a comprehensive, authoritative compendium of human genes and genetic phenotypes that is freely available and updated daily. The full-text, referenced overviews in OMIM contain information on all known mendelian disorders and over 12,000 genes. OMIM focuses on the relationship between phenotype and the entries contain copious links to other genetics resources.

- **RareConnect** ([https://www.rareconnect.org/en](https://www.rareconnect.org/en))
  RareConnect was created by EURORDIS (European Rare Disease Organisation) and NORD (National Organization for Rare Disorders) to provide a safe space where individuals and families affected by rare diseases can connect with each other, share vital experiences, and find helpful information and resources

- **Wolfram Syndrome UK**: [www.wolframsyndrome.co.uk](http://www.wolframsyndrome.co.uk)
  The website is run by families affected by this rare genetic disorder and the aim is to raise as much awareness of the syndrome as possible.