

Child and Adolescent Mental Health Services (CAMHS)

Effective shared care agreement (ESCA) for the treatment of Attention Deficit Hyperactivity Disorder (ADHD) with DEXAMFETAMINE

This form must be completed by the hospital specialist/consultant (may include specialist nurse non-medical prescribers) and sent to the GP for approval. It must be signed by the GP and returned to the team before the patient is informed that the GP will prescribe the medication.

SECONDARY CARE SECTION TO BE COMPLETED BY INITIATING CONSULTANT/SPECIALIST

Patient's Name: _____	NHS Number: _____
Date of Birth: _____	ESCA Date: _____
One copy of information leaflet given to patient	
One copy of agreement sent to GP	
One copy filed in patients notes	
Consultant/Specialist Name (please print): _____	
Consultant/Specialist Signature: _____	
Support contact number: _____ (if not listed overleaf)	
Email address / Fax no: _____	

PRIMARY CARE SECTION TO BE COMPLETED BY GENERAL PRACTITIONER

I agree*/don't agree* to enter into a shared care arrangement for the treatment of the above patient with this medicine (*delete as appropriate)	
GP Name: _____	
Signature: _____	Date: _____
Once signed please email or fax back to the team.	

CONSENT SECTION TO BE COMPLETED BY PATIENT / REPRESENTATIVE

I agree*/don't agree* to enter into a shared care arrangement for the above treatment (*delete as appropriate)

Patient / Representative Name: _____ **Signature:** _____

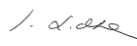


Date: _____

BACK UP ADVICE AND SUPPORT

Contact details	Telephone number
Sandwell Base	0121 612 6620

Contact details	Telephone number
Wolverhampton Base	01902 444 021

Version Control			
Version	Date of Approval	Author/s	Brief Description of Changes
1.0	2006	Dr Win	New ESCA
2.0	July 2013	Dr Win	Revised front page, new signatories
4.0	Nov 2018	Mr Narinder Sangha	Separated out individual drugs for treatment of ADHD into separate ESCA's. Now includes hospital specialists/ NMP's and parents/carers.

This shared care agreement has been approved for use by:		Signature	Date
BCPFT MMC Chair	Dr J Lidher		15/10/2018
Wolverhampton City CCG Prescribing Lead	Dr A Stone		18/12/2018
Sandwell & West Birmingham CCG Medicines Quality	Jonathan Boyd		20 th Dec 2018

AREAS OF RESPONSIBILITY FOR THE SHARING OF CARE

The aim of an Effective Shared Care Agreement (ESCA) is to provide information to general practitioners (GPs) and hospital staff about complex or high cost therapies that their patients may receive following specialist referral. An ESCA will be written only when it has been agreed that shared care is an appropriate option, and will include a statement of specialist and GP responsibilities.

ESCA's will ensure that all GPs have sufficient information to enable them to undertake prescribing responsibility for specialist therapies and other therapies that may affect or interact with specialist therapies.

It is not the intention to insist that GPs prescribe this therapy and any doctor who does not wish to undertake the clinical and legal responsibility for this drug is not so obliged.

The doctor who prescribes the medication legally assumes clinical responsibility for the drug and the consequences of its use.

RESPONSIBILITIES AND ROLES

Specialist responsibilities

- Arrange comprehensive assessment of the child and be responsible for making the diagnosis and considering alternative diagnoses, co-morbid diagnoses and cautions/ contraindications to treatment.
- Initiate treatment with stimulants. Prescribe by brand for sustained release preparations. Inform the GP promptly about changes in treatment or dosage, any important adverse events or if other interacting medicines are prescribed/recommended.
- Monitor the patient's condition and response to treatment regularly and keep the GP informed.
- Provide a comprehensive baseline physical assessment as recommended in the NICE guidelines, and ensure that height, weight, blood pressure, pulse and appetite are monitored at the recommended time intervals using the chart in the appendix. Communicate the results of tests to the GP as soon as possible.
- Explain the possible side effects of the drug and interactions to parents.
- Provide written guidance for parents (at specialist's discretion).
- Be available for back-up advice on any of the above during working hours.
- Report all suspected adverse drug reactions (in children under 18 years of age), to the Medicines and Healthcare Regulatory Agency (MHRA) via www.mhra.gov.uk/yellowcard
- Send a letter and shared care agreement form to the GP to obtain consent to share prescribing and monitoring responsibilities.
- Advise GP's of any dosage adjustments required, when to refer back, and when and how to stop treatment.
- Ensure clear arrangements for back up, advice and support.

General Practitioner responsibilities

- Prescribe **Dexamfetamine** once notified by the specialist.
- Ensure that the treatment is not continued if the patient fails to attend the specialist clinic for over a year.
- Check that patient is being monitored as specified in section on specialist responsibility.
- Report to and seek advice from the specialist on any aspect of patient care that is of concern to the GP and may affect treatment.
- Refer back to the specialist if the patient's condition deteriorates.
- Monitor the patient for side effects and report all suspected adverse drug reactions (in children under 18 years of age), to the specialist and to the MHRA via www.mhra.gov.uk/yellowcard
- Stop the treatment if advised by the specialist.

Parent/carers role

- Ask the specialist or GP anything he or she does not understand about the treatment.
- Try to put into practice any behavioural or psychological programmes and report back to the specialist about their effectiveness.
- Report any adverse effects to the specialist or GP.
- Attend agreed review appointments.

SUPPORTING INFORMATION

Licensed indications

Attention Deficit Hyperactivity Disorder (ADHD) is a heterogeneous behavioural syndrome characterised by the core symptoms of inattention, hyperactivity and impulsivity. ADHD should only be diagnosed by a specialist psychiatrist, paediatrician, or other healthcare professional with training and expertise in the diagnosis of ADHD. Diagnosis of ADHD should be made according to DSM-IV criteria or the guidelines in ICD-10 and should also be based on a complete history and evaluation of the patient, including a full clinical and psychosocial assessment, full developmental and psychiatric history, and assessment of mental state. Diagnosis cannot be made solely on the presence of one or more symptom.

Dexamfetamine is licensed for the treatment of ADHD in the UK. It should form part of a comprehensive treatment programme for ADHD that includes psychological, behavioural and educational advice and interventions. The indication for drug treatment is the presence of impairment resulting from ADHD.

Adverse events should be reported. Reporting forms and information can be found at www.mhra.gov.uk/yellowcard

Therapeutic Use

The aim of stimulant medication as part of a comprehensive treatment programme is to stabilise children with a behavioural syndrome characterised by symptoms which may include chronic history of short attention span, distractibility, emotional lability, impulsivity and moderate to severe hyperactivity.

DEXAMFETAMINE

Dexamfetamine is used in children with refractory ADHD who do not respond to Methylphenidate or Atomoxetine.

Dosage and Administration

Child 6-17 years: Initially 2.5 mg – 2-3 times daily, increased if necessary at weekly intervals by 5mg, usual max.1mg/kg daily, up to 20 mg daily (40 mg daily has been required in some children).

Formulation

Tablets may be halved.

Contraindications

Cardiovascular disease including moderate to severe hypertension, structural cardiac abnormalities, advanced arteriosclerosis, hyper-excitability or agitated states, hyperthyroidism, history of drug or alcohol abuse.

Cautions

Mild hypertension, anorexia, psychosis or bipolar disorder, monitor for aggression and hostility during initial treatment, history of epilepsy, tics and Tourette syndrome, monitor growth in children, susceptibility to angle-closure glaucoma, avoid abrupt withdrawal, acute porphyria.

Side effects

Common or very common side effects include; abdominal cramps; acidosis; aggression; alopecia; anhedonia; anorexia; anxiety; ataxia; cardiomyopathy; cardiovascular collapse; cerebral vasculitis; chest pain; confusion; depression; diarrhoea; dizziness; dry mouth; dysphoria; euphoria; growth restriction in children; headache; hyperactivity; hyperpyrexia; hyperreflexia; hypertension; hypotension; impaired concentration; irritability; ischaemic colitis; malaise; mydriasis; myocardial infarction; nausea; nervousness; neuroleptic malignant syndrome; obsessive-compulsive behaviour; palpitations; panic attack; paranoia; psychosis; rash; renal impairment; restlessness; rhabdomyolysis; seizures; sexual dysfunction; sleep disturbances; stroke; sweating; tachycardia; taste disturbance; Tourette syndrome (in predisposed individuals); tremor; urticaria; visual disturbances; weight loss.

Monitoring

Monitoring (see appendix for monitoring chart)	Pre-treatment assessment	Prior to prescribing, it is necessary to conduct a baseline evaluation of a patient's cardiovascular status including blood pressure and heart rate. A comprehensive history should document concomitant medications, past and present co-morbid medical and psychiatric disorders or symptoms, family history of sudden cardiac/unexplained death and accurate recording of pre-treatment height and weight on a growth chart.		
	After commencing treatment	Growth, psychiatric and cardiovascular status should be continuously monitored. <ul style="list-style-type: none"> • Blood pressure and pulse should be recorded on a chart at each adjustment of dose and then at least every 6 months; • Height, weight and appetite should be recorded at least 6 monthly with maintenance of a growth chart; • Development of de novo or worsening of pre-existing psychiatric disorders, including depression and aggressive behaviour, should be monitored at every adjustment of dose and then at least every 6 months and at every visit. Patients should be monitored for the risk of diversion, misuse, and abuse of Dexamfetamine .		
	Actions to be taken:			
	Parameter	Monitoring frequency	Action	By Whom
	Weight gain / Appetite	At least 6 monthly	Failure to gain weight appropriately - may require withdrawal.	Specialist at regular reviews.
Blood pressure / pulse	At least 6 monthly	Monitor whilst taking medication to ensure within published range for age of child.		
Growth Development	At least 6 monthly	If adversely affected consideration should be given to dose reduction or interrupting therapy in those on long-term treatment.		

NICE CG72: Drug holidays are not routinely recommended; however, consideration should be given to the parent or carer and child or young person with ADHD working with their healthcare professional to find the best pattern of use, which may include periods without drug treatment. If growth is significantly affected by drug treatment, the option of a planned break in treatment over school holidays should be considered to allow 'catch-up' growth to occur.

Treatment cessation

Abrupt withdrawal should be avoided.

For full details of the licensed indications, dosage and administration, cautions, contraindications, side effects, drug interactions and monitoring, refer to the current edition of the BNF, BNFc and SPC.

Dexamfetamine is a **Schedule 2 Controlled Drug**. Legislation for controlled drug prescription writing is set out in the BNF in the section on 'Controlled Drugs and Drug dependence'.

References

- British National Formulary. BMJ Group/Royal Pharmaceutical Society of Great Britain. BNF App Accessed August 2017.
- Summary of Product Characteristics (SPC). Accessed October 2017 via www.medicines.org.uk/emc
- National Institute for Health and Clinical Excellence. Clinical Guideline 72. *Attention deficit hyperactivity disorder: Diagnosis and management*. September 2008. Last updated February 2016.
- Taylor D, Paton C, Kapur S. The Maudsley Prescribing Guidelines in Psychiatry. Twelfth edition. The South London and Maudsley NHS Foundation Trust, Oxleas NHS Foundation Trust. 2015.
- Dexamfetamine ESCA (as Amfexa). Birmingham, Sandwell and environs Area prescribing Committee (BSSE APC) 2017.

Appendix

Chart for ongoing monitoring during Dexamfetamine Sulfate treatment For specialist to complete

As outlined in the prescribing information in more detail, growth, psychiatric and cardiovascular status should be regularly monitored:

- Blood pressure and pulse should be recorded at each adjustment of dose and then at least every 6 months
- Height, weight and appetite should be recorded at least 6-monthly with maintenance of a growth chart
- Development of *de novo* or worsening of pre-existing psychiatric disorders should be monitored at every adjustment of dose and then at least every 6 months and at every visit

Date of initial assessment: _____ Patient name: _____

Date of birth: _____ NHS No: _____

	Baseline	Subsequent appointments					
Date of assessment							
Reason for assessment							
Blood pressure*							
Heart rate*							
Body weight (kg)**							
Height (cm)**							
Appetite							

*Blood pressure and heart rate should be recorded at each adjustment of dose and then at least every 6 months

**Height, weight and appetite should be recorded at least 6-monthly with maintenance of a growth chart