SHARED CARE PRESCRIBING GUIDELINE

Methylphenidate, atomoxetine, lisdexamfetamine, dexamfetamine and guanfacine for the treatment of Attention Deficit Hyperactivity Disorder in Children and Adolescents aged 6-18 years

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| **NOTES to the GP** |
| The information in the shared care guideline has been developed in consultation with South East London ICB and it has been **agreed** that it is suitable for shared care.This document should provide sufficient information to enable you to make an informed decision regarding the clinical and legal responsibility for prescribing either **methylphenidate, atomoxetine, lisdexamfetamine, dexamfetamine or guanfacine for the treatment of ADHD in Children and Adolescents aged 6-18 years\***The questions below will help you confirm this: * Is the patient’s condition predictable or stable?
* Do you have the relevant knowledge, skills and access to equipment to allow you to monitor treatment as indicated in this shared care prescribing guideline?
* Have you been provided with relevant clinical details including monitoring data?

**If you can answer YES to all these questions** (after reading this shared care guideline), then it is appropriate for you to accept prescribing responsibility.**If the answer is NO to any of these questions** you should contact the requesting specialist or your borough Medicines Optimisation team. There may be implications for the patient where the invitation to share care is declined. For example, the patient may need to be changed to an alternative treatment regimen. **It would not normally be expected that shared care prescribing would be declined on the basis of cost.** Sharing of care assumes communication between the specialist, GP and patient. The intention to share care should be explained to the patient by the specialist initiating treatment. **It is important that patients are consulted about treatment and are in agreement with it**. Prescribing should follow requirements in the South East London Interface Prescribing Policy. **The doctor who prescribes the medication legally assumes clinical responsibility for the drug and the consequences of its use. The patient’s best interests are always paramount.****\*N.B** *NICE recommends that the treatment of ADHD can start from the age of 5 years – however all medicines that are used to treat ADHD are only licensed for children from 6 years of age.*  |

**Once you have read the shared care guideline and considered the information above, please complete the GP decision form on the next page and email back to the requesting clinician if you are in agreement to participate in shared care. If you are not in agreement, please include reasons for this.**

**GP DECISION FORM**

This shared care agreement outlines suggested ways in which the responsibilities for managing the prescribing of **methylphenidate, atomoxetine, lisdexamfetamine, dexamfetamine and guanfacine for the treatment of ADHD** in **children and adolescents aged 6-18 years** can be shared between the specialist and general practitioner (GP). GPs are invited to participate. If the GP is not confident to undertake these roles, then he or she is under no obligation to do so. In such an event, the total clinical responsibility for the patient for the diagnosed condition remains with the specialist. If a specialist asks the GP to prescribe this drug, the GP should reply to this request as soon as practicable.

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| **AGREEMENT TO PARTICIPATE IN SHARED CARE****Of [***either* ***methylphenidate or atomoxetine or lisdexamfetamine* *or dexamfetamine or guanfacine*** *delete and enter drug name above as appropriate***]** **for the treatment of ADHD in Children and Adolescents aged 6-18 years** |
| **Consultant/Specialist Name:** | **Patient name:** |
| **Consultant/Specialist signature:** | **Patient Hospital Number:****Patient NHS Number:**  |
| **Date completed:** | **Patient/Carer Agreement:**Patient/Carer agrees to shared care □Patient/Carer does not agree to shared care □ |
| **Hospital requesting shared care:** |
| **GP Name:** |
| **This is to confirm that I agree/do not agree [delete as appropriate] to participate in shared care for (***Enter drug name*…………………....) **for the treatment of ADHD** **for this patient as outlined in this shared care document** |
| **GP Signature:** | **Date signed:**  |
| If **NOT** happy to take on shared care prescribing of this medication – please state why: |
| **ACTION**1. **HOSPITAL CONSULTANT/SPECIALIST Tick to confirm**

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| * Explain shared care to patient/carer and obtain agreement. Date of agreement:………...……..
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| * Indicate requesting hospital
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| * Complete and sign agreement
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| * Email (via secure email) full shared care guideline (including signed agreement to GP,
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| * Place original in patient’s notes
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1. **GP PRACTICE**
* If **in agreement** to participate in shared care, sign and email (via secure NHS.net) this sheet back **within 2 weeks** of receipt **of request from specialist** to either:

Email address (via secure nhs.net):……………………………………. **[TRUST to ADD email address] OR*** Ifyou **do not agree** to participate in shared care, contact consultant/specialist and borough Medicines Optimisation Team within 2 weeks of receipt to discuss. If after discussion it is agreed not to undertake shared care for this patient, both the consultant/specialist and the borough Medicines Optimisation team should be informed.
* Once a decision is reached, file a copy in the Patient’s medical notes.
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**Methylphenidate, atomoxetine, lisdexamfetamine, dexamfetamine and guanfacine for the treatment of ADHD in** **Children and Adolescents aged 6-18 years**

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| **Drug Name** | **Licensed Indication** | **Preparations** |
| **METHYLPHENIDATE****(CD Schedule 2)** **Prescriptions for sustained release tablets or capsules should specify the brand** | **Licensed for ADHD for children over 6 years of age. First line for ADHD** | ***Plain tablets*:** **Available in the following strengths:** **5mg, 10mg, 20mg***Brands include Ritalin® Medikinet®*N.B. any equivalent strength tablet can be prescribed ***Sustained Release TABLETS (Xenidate****®* ***XL, Delmosart****®* ***prolonged-release tablets and Concerta****®* ***XL)*:** **Available in the following strengths:** **18mg, 27mg, 36mg, 54mg** *NB – These three brands are bioequivalent. Xenidate® XL or Delmosart® prolonged-release tablets should be prescribed first-line. Concerta® XL tablets remain as a third-line option if patient’s ADHD control destabilizes on Xenidate® XL and Delmosart® prolonged-release tablets).****Sustained Release CAPSULES:*** **Available in the following strengths:** **5mg, 10mg, 20mg, 30mg, 40mg, 50mg, 60mg***Brands include**Medikinet® XL**Equasym® XL Meflynate® XL*N.B. any equivalent strength sustained release capsule can be prescribed but patients should remain on the same brand that they are initiated on |
| **ATOMOXETINE** | **Licensed for ADHD for children over 6 years of age.** | **Strattera®capsules 10mg, 18mg, 25mg, 40mg, 60mg, 80mg, 100mg** **Oral solution 4mg/ml** |
| **LISDEXAMFETAMINE****(dimesylate)****(CD Schedule 2)** | **Licensed for ADHD for children over 6 years of age.** | **Elvanse® capsules 20mg, 30mg, 40mg, 50mg,60mg and 70mg**  |
| **DEXAMFETAMINE****(sulphate)****(CD Schedule 2)** | **Licensed for ADHD for children over 6 years of age.** | **Amfexa® tablets 5mg, 10mg 20mg** **Oral solution 1mg/ml** |
| **GUANFACINE** | **Licensed for ADHD for children and adolescents 6-17 years old** | **Intuniv® tablets 1mg, 2mg, 3mg 4mg** |

# **CIRCUMSTANCES WHEN SHARED CARE IS APPROPRIATE**

* Prescribing responsibility will only be transferred when the consultant/specialist and the GP are in agreement that the patient’s condition is stable or predictable.
* On initiation of treatment the consultant/specialist will provide prescriptions for a minimum of 12 weeks (if CD schedule 2 drug supply either as 3x28 or 3x30 day prescriptions depending on pack size)
1. **Areas of responsibility**

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| **Consultant / Specialist team responsibilities** |
| ***Before requesting agreement for shared care**** Establish or confirm diagnosis and assess patient suitability for treatment
* Conduct a careful history and physical examination to assess any presence of cardiac disease
* Establish and document any allergies and previous hypersensitivity

Baseline monitoring - **These should be shared with the GP following a request to take up shared care*** Height and weight – add to a growth chart
* Cardiovascular status, including blood pressure and heart rate before prescribing and obtain specialist cardiac advice if appropriate
* Discuss treatment with patients or carers, ensuring and documenting that they have a clear understanding of benefits, side effects, frequency of administration and monitoring requirements
* Email a signed shared care guideline with patient details completed to GP for consideration of shared care request
* Before treatment is initiated check for any potential drug interactions if patient is currently on other medications
* Initiate treatment and titrate the dose against symptoms and side effects over 4-6 weeks until dose optimisation is achieved.
* Prescribe and monitor treatment according to local guideline or formulary until patient’s condition is stable or predictable
* At the time of initiating, inform GP in writing as to which of the 5 drugs included in this shared care guideline has been prescribed and **to clarify this on page 2 of this agreement**.
* The GP should be invited to share care once the patient is stable. Information provided to the GP should include:
* A copy of the shared care guidelines with the relevant amendments made under *Agreement to participate in shared care* (page 2) detailing the drug which will involve shared care.
* That prescriptions for a minimum of 12 weeks supply has or will be given
* Information on when the patient will next be reviewed and by whom.
* A request that the GP continue prescribing after 12 weeks.
* Advise GP on the appropriateness of any necessary periodic drug holidays

***After agreement to shared care**** Inform GP when patient is stable see above – dose titration should occur before transfer.
* Inform GP of abnormal monitoring results and any changes in therapy
* Evaluate adverse events reported by GP or patient
* Carry out ongoing monitoring and follow up according to shared care guidelines including continued need for therapy.
* If a dose change is needed, a prescription is issued from the clinic and GP provided with a letter of the dose change and information regarding any further monitoring that may be required. Consultant/Specialist should review the patient within 3-6 months following any dose change. Advise GP when ADHD treatment should be discontinued and provide necessary supervision and support during the discontinuation phase.
* To communicate promptly with the GP if treatment is changed.
* To report any suspected adverse effects to the MHRA: <http://www.yellowcard.gov.uk>
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| **General Practitioner responsibilities**  |
| ***Before agreement to shared care**** Consider shared care proposal within 2 weeks of receipt and fill in GP Decision form (page 2) and return to specialist.
* State in the patient’s records that the medicine is being prescribed under a shared care agreement

***After agreement to shared care**** Prescribe dose as recommended once the patient’s condition is stable or predictable.
* Add ‘shared care’ read code to patient’s medical record.
* Continue prescriptions after stabilisation in line with the points below.
* Monitor general health of patient and check adverse effects as appropriate
* Monitor height, weight, (check against <https://www.rcpch.ac.uk/resources/growth-charts>) blood pressure\* and pulse after the first 3 months of treatment, as well as after each dose adjustment as directed by the specialist, and then every 6 months. Any significant changes from baseline in BP/weight/pulse should be discussed with the specialist.. If patients develop symptoms suggestive of cardiac disease during treatment, they should be referred for prompt specialist cardiac evaluation and the consultant/specialist team informed
* Stop treatment on advice of specialist or immediately if urgent need arises
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| **General Practitioner responsibilities (continued)**  |
| * Check for drug interactions when prescribing new or stopping existing medication
* Discuss any abnormal results with specialist and agree any action required (this could be a telephone discussion).
* Only ask specialist to take back prescribing should the patients clinical condition deteriorate. Allow an adequate notice period of 10 working days. Consider a telephone discussion with the specialist if appropriate.
* Check that the patient is attending specialist appointments at least annually
* To advise the specialist if non-compliance is suspected
* To report any suspected adverse effects to the MHRA via the Yellow Card scheme: <http://www.yellowcard.gov.uk>

\*GP’s can order small cuffs to enable them to monitor blood pressure, where they are unable to carry out this monitoring they should inform the specialist to arrange how blood pressure can be monitored. |
| **Patient's / Carer’s responsibilities** |
| * To contact the specialist or GP if he or she does not have a clear understanding of any aspect of the treatment.
* To inform prescribing specialist, GP and other healthcare professionals of any other medication being taken, including over the counter products, alternative therapies or recreational drugs.
* To inform community pharmacists that they are using ADHD Treatments before purchasing medication over-the-counter
* To attend all hospital and GP appointments
* To take medicines as agreed and take steps to ensure that no doses are missed and not to share medicines with others
* To read the patient information leaflet included with the medication.
* To report any adverse effects or warning symptoms to GP or hospital specialist
* To report to GP if pregnant or breastfeeding.
* To inform GP and hospital of any changes in addresses or telephone contact numbers.
* To request the need for repeat prescriptions in a timely manner to allow appropriate processing of the script. **N.B**. If patient is prescribed methylphenidate, dexamfetamine or lisdexamfetamine these prescriptions will be issued as paper prescriptions and be picked up from the GP and taken to local pharmacy for dispensing
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1. **CLINICAL INFORMATION**

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| **M METHYLPHENIDATE, ATOMOXETINE LISDEXAMFETAMINE, DEXAMFETAMINE AND****GUANFACINE****Monitoring Requirements including frequency** |
| **Consultant/Specialist:** * To assess baseline cardiovascular status, including blood pressure and heart rate before prescribing and obtain specialist cardiac advice if appropriate.
* To review the patient and monitor the following on an annual basis for the duration that the patient is on the medicine and communicate these results to the GP:
* For children under 10 years monitor height, weight and appetite, recorded on a growth chart (check against <https://www.rcpch.ac.uk/resources/growth-charts>)
* Blood pressure and pulse, recorded on a centile chart (also following dose adjustments).
* To refer patients who develop symptoms such as palpitations, exertional chest pain, unexplained syncope, dyspnoea, or other symptoms suggestive of heart disease for prompt specialist cardiac evaluation.
* The development of new or worsening of pre-existing, psychiatric symptoms (also following dose adjustments and at every visit).
* Monitoring of motor / verbal tics should be carried out at every dose adjustment and at least annually.
* Blood testing should be carried out periodically at the discretion of the supervising clinicians and when clinically indicated (e.g. if recurrent nose bleeds, bruising or infections occur).

*Methylphenidate, dexamfetamine and lisdexamfetamine are classed as* ***controlled drugs (see page 7) for prescribing information)****,* ***Atomoxetine and Guanfacine are Prescription Only Medicines.*** *In order to monitor the effects of treatment the specialist or parents should inform the school concerning any medication for these indications. In order to assess the effects of the drug on the child’s emotional, physical or behavioral states the specialist should request further information from the school about the child’s behaviour.*  |
| **M METHYLPHENIDATE, ATOMOXETINE LISDEXAMFETAMINE, DEXAMFETAMINE AND****GUANFACINE****Monitoring Requirements including frequency (continued)** |
| **GP:*** To monitor, pulse, blood pressure\*, and height and weight (for children under 10 years old - check against <https://www.rcpch.ac.uk/resources/growth-charts>) every three months
* To contact specialist if deterioration in behaviour.
* To report adverse drug reactions to specialist.
* To refer patients who develop symptoms such as palpitations, exertional chest pain, unexplained syncope, dyspnoea, or other symptoms suggestive of heart disease for prompt specialist cardiac evaluation.
* To refer patients with recurrent nose bleeds, bruising or infection.

Blood pressure and pulse rate checks should ideally be done in the more relaxed environment of a GP surgery rather than in hospital. But in reality, the BP and pulse should be checked by whoever sees the patient first after a dose increase (usually within 2 weeks of the change)\*GP’s can order small cuffs to enable them to monitor blood pressure, where they are unable to carry out this monitoring they should inform the specialist to arrange how blood pressure can be monitored |
| **Follow up arrangements** |
| **Consultant/Specialist:*** To arrange for follow up at least annually and following each dose adjustment
* Arrangement of a clinic review when the patient is between 17 to 18 years should be considered to assess continued treatment into adult services and to plan for the transfer of care if needed

**GP:*** To act upon results communicated by specialist.
* To review the appropriateness of prescribing for patients who have not been seen by a specialist for over one year.
* Communicate with the consultant/specialist if the patient does not attend appointments
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| **Duration of treatment**  |
| Long-term treatment may continue into adulthood. Patients who take treatment for extended periods (i.e. >1 year) should have their treatment reviewed at least once a year by a specialist to determine whether continuation is needed |
| **Criteria for stopping treatment** |
| If improvement of symptoms is not observed after the appropriate dosage adjustment over one month, it should be discontinued. The drug may be discontinued periodically (e.g. by stopping the drug for up to two weeks each year) to assess the child’s condition as advised by the consultant/specialist. Need for continued treatment should be routinely reviewed beyond the age of 18 years |

***NOTE:*** *The Information here is* ***not*** *exhaustive.* ***Please consult the current Summary of Product Characteristics (SPC) for the treatment prior to prescribing for up to date prescribing information including detailed information on adverse effects, drug interactions, cautions and contraindications (available via*** [***www.medicines.org.uk***](http://www.medcines.org.uk/)***)***

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| **Drug** | **Indication** | **Place in therapy** | **Dose and Route of Administration** |
| **Preparation** | **Dose** | **Notes** |
| Methylphenidate | Treatment of ADHD | First line for ADHD | **Plain tablets\***Available in the following strengths: **5mg, 10mg, 20mg**Ritalin® Medikinet®  | Initially 5 mg 1–2 times a day, increased in steps of 5–10 mg daily if required, at weekly intervals, increased if necessary up to 2.1 mg/kg daily in 2–3 divided doses, max. licensed dose is 60 mg daily in 2–3 doses, (maximum of 90 mg daily under the direction of a specialist) discontinue if no response after 6 weeks | In some children rebound hyperactivity may occur if the effect of the drug wears off in the evening. An additional dose later in the day may eliminate this difficulty but may disturb sleep.  |
| **Sustained release tablets**Available in the following strengths **18mg, 27mg, 36mg, 54mg** Xenidate® XL, Delmosart® prolonged-release tablets, Concerta® XL***The prescriber must specify the brand***  | Initially 18 mg once daily in the morning, increased in steps of 18 mg daily at weekly intervals, increased if necessary up to 2.1 mg/kg daily, max. licensed dose is 54 mg daily, (maximum of 108 mg daily under the direction of a specialist) discontinue if no response after 6 weeks | Total daily dose of 15mg of standard release tablet is considered equivalent to18mg once daily of sustained release tablets. 60mg of Ritalinis the maximum licensed dose. The equivalent dose of Concerta**®** XLis 72mg, which is above the maximum licensed dose.  |
| **Sustained release capsules**Available in the following strengths **5mg, 10mg, 20mg, 30mg, 40mg, 50mg,60mg**Equasym® XL Medikinet® XL Meflynate® XL***The prescriber must specify the brand*** | Initially 10mg once daily (in the morning before breakfast), increasing if necessary, by weekly increments of 10mg to a max. licensed dose of 60 mg daily, (maximum of 90 mg daily under the direction of a specialist) discontinue if no response after 6 weeks | 40mg XL strength not available in Equasym® XLbrandMeflynate XL® may be taken independent of food intake; Equasym XL® should be taken before food; Medikinet XL® should be taken with or after food.  |
| Lisdexamfetamine | Licensed for ADHD for children over 6 years of age. | To be considered if methylphenidate has not been successful or tolerated | **Elvanse® 20mg. 30mg, 40mg, 50mg 60mg and 70mg Capsules** | Starting dose 30mg taken once in the morning (with or without food)The dose may be increased by10-20mg increments at approximately weekly intervals.Maximum recommended dose = 70mg/day | Lower starting dose of 20mg once daily may be needed in some patientsLisdexamfetamine may be swallowed whole, or the capsules opened and the entire contents emptied and mixed with a soft food such as yogurt or in a glass of water or orange juice |
| Dexamfetamine | Licensed for ADHD for children over 6 years of age. | To be considered if methylphenidate not successful or tolerated and have responded to lisdexamfetamine but cannot tolerate the longer effect profile | **Amfexa® tablets 5mg, 10mg 20mg**Oral solution is 1mg/ml  | Initially 2.5 mg 2–3 times a day, increased in steps of 5 mg once weekly if required, increased if necessary up to 1 mg/kg daily, maintenance dose to be given in 2–4 divided doses, up to 20 mg daily (40 mg daily has been required in some children). |  |
| **Drug** | **Indication** | **Place in therapy** | **Dose and Route of Administration** |
| **Preparation** | **Dose** | **Notes** |
| Atomoxetine | Licensed for ADHD for children over 6 years of age. | To be considered if methylphenidate or lisdexamfetamine has not been successful or tolerated | **Strattera® Capsules 10mg, 18mg, 25mg, 40mg, 60mg, oral solution 4mg/ml****Child over 6 years (body-weight <70kg)** | Initially 500 micrograms/kg daily for 7 days, increased according to response; usual maintenance dose 1.2mg/kg daily, but may be increased to 1.8mg/kg daily (max. 120mg daily) under the direction of a specialist | The SPC dosing states that: *“No additional benefit has been demonstrated for doses higher than 1.2mg/kg/day. The safety of single doses over 1.8mg/kg/day and total daily doses above 1.8mg/kg has not been systematically evaluated.”* 4 The 1.2mg/kg/day dose is based on 2001 data on uncomplicated ‘pure’ attention deficit hyperactivity disorder. The consultant-led clinic is full of complex patients with co-morbidities, and since 2005 a 1.8mg/kg/day dose is known to be more effective in this group4. |
| **Child over 6 years (body-weight >70kg)** | Initially 40mg daily for 7 days, increased according to response; usual maintenance dose 80mg daily, but may be increased to 120mg daily under the direction of a specialist. |
| Guanfacine  | Licensed for ADHD for children over 6 years of age | To be considered if methylphenidate or lisdexamfetamine has not been successful or tolerated | **Intuniv ® tablets 1mg, 2mg, 3mg, 4mg** **Child 6-17years (body-weight 25kg – 41.4kg)** | Initially 1mg once daily increasing in weekly increments of 1mg up to a maximum of 4mg once daily  |  |
| **Child 13 - 17years (body-weight 41.5kg – 49.4kg)** | Initially 1mg once daily increasing in weekly increments of 1mg up to a maximum of 5mg once daily |  |
| **Child 13 - 17years (body-weight 49.5kg – 58.4.kg)** | Initially 1mg once daily increasing in weekly increments of 1mg up to a maximum of 6mg once daily |  |
| **Child 13 - 17years (body-weight >58.4.kg)** | Initially 1mg once daily increasing in weekly increments of 1mg up to a maximum of 7mg once daily | Dose can be titrated to a 7 mg/day dose after the subject has completed a minimum of 1 week of therapy on a 6 mg/day dose and the physician has performed a thorough review of the subject's tolerability and efficacy. |

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| **Further Information**  |
| A pharmaceutical company patient information leaflet (PIL) will be provided to the patient with each supply. Medicines for children have also produced PIL which can be accessed via the online website<http://www.medicinesforchildren.org.uk/search-for-a-leaflet/> NICE has produced an information leaflet for parents:<http://www.nice.org.uk/nicemedia/pdf/CG72UNG.pdf> A review letter will be sent after initial assessment and following each further appointment. It is assumed that the GP agrees to the shared care arrangements.**Information which can be provided to the schools -** Managing Medicines in Schools and Early Years Settings<https://www.education.gov.uk/publications/standard/publicationdetail/page1/DFES-1448-2005>**Information on prescribing Controlled Drugs** Methylphenidate, lisdexamfetamine and dexamfetamine are schedule 2 Controlled drugs - the following applies:* Prescribers can now issue computer-generated prescriptions for all CDs including Schedule 2 and 3 CDs; all details except the signature can be computer-generated
* Prescriptions for Schedule 2 CDs are only valid for 28 days.
* Schedule 2 CDs cannot be prescribed on repeat dispensing prescriptions
* There is a good practice requirement that the quantity of Schedule 2 CDs be limited to a quantity for up to 30 days treatment. In cases where the prescriber believes that a prescription should be issued for a longer period they may do so but will need to be able to justify that there is a clinical need and that it would not cause an unacceptable risk to patient safety
* The prescription for CDs must contain the dose, form, strength (where appropriate) and a total quantity of the preparation in both words and figures
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| **References** |
| NICE. Clinical Guideline 87: Attention deficit hyperactivity disorder: Diagnosis and Management (March 2018). Accessed via: <https://www.nice.org.uk/guidance/ng87>NICE Technology Appraisal Number 98 Methylphenidate, atomoxetine and dexamfetamine for attention deficit hyperactivity disorder (ADHD) in children and adolescents. March 2006 www.nice.org.uk NICE. Clinical Guideline 72: Attention deficit hyperactivity disorder: Diagnosis and management of ADHD in children, young people and adults (2008). Accessed via [this link](http://publications.nice.org.uk/attention-deficit-hyperactivity-disorder-cg72) (superseded by NG87)British National Formulary for Children 2016/17**Summary of Product Characteristics** – accessed via [www.medicines.org.uk](http://www.medicines.org.uk) 1. Ritalin® - (Last accessed April 2018)
2. Equasym® - (Last accessed April 2018)
3. Equasym XL® - (Last accessed April 2018)
4. Medikinet® - (Last accessed April 2018)
5. Medikinet XL® - (Last accessed April 2018)
6. Concerta XL® - (Last accessed April 2018)
7. Strattera® - (Last accessed April 2018)
8. Elvanse® - (Last accessed April 2018)
9. Xenidate®XL (Last accessed April 2018)
10. Amfexa ® (Last accessed April 2018)
11. Intuniv® (Last accessed April 2018)
12. Delmosart® (Last accessed April 2018)
13. Meflynate ® (Last accessed March 2024)
14. Atomoxetine Treatment in Children and Adolescents with Attention-Deficit/Hyperactivity Disorder and Comorbid Oppositional Defiant Disorder. Newcorn J H et al. Journal of the American Academy of Child and Adolescent Psychiatry. 1 March 2005 (vol 44 issue 3 pages 240-8)

NICE ESNM19: Attention deficit hyperactivity disorder in children and young people: lisdexamfetamine dimesylate (May 2013). Accessed via [this link](http://publications.nice.org.uk/esnm19-attention-deficit-hyperactivity-disorder-in-children-and-young-people-lisdexamfetamine-esnm19) 1. Scottish Medicines Consortium. Lisdexamfetamine dimesylate (May 2013). Accessed via <http://www.scottishmedicines.org.uk/files/advice/lisdexamfetamine_dimesylate__Elvanse__FINAL_April_2013_Amended_26.04.13_for_website.pdf>
2. Coghill D, Banaschewski T et al. [European, randomized, phase 3 study of lisdexamfetamine dimesylate in children and adolescents with attention-deficit/hyperactivity disorder.](http://dx.doi.org/10.1016/j.euroneuro.2012.11.012) European Neuropsychopharmacology 2013; doi:10.1016/j.euroneuro.2012.11.012
3. Dittmann RW, Cardo E et al. Efficacy and safety of lisdexamfetamine dimesylate and atomoxetine in the treatment of Attention-Deficit/Hyperactivity Disorder: a head-to-head, randomised, double blind, Phase IIIb study. CNS Drugs 2013 DOI 10.1007/s40263-013-0104-8
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# **COMMUNICATION AND SUPPORT**

# **Please note that the clinical letter received from the consultant/specialist team should have the relevant contact details. If this is not provided you may find the following contact details useful.**

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| **King’s College (Denmark Hill and Princess Royal Hospital Sites) switchboard: 020 3299 9000** |
| **Medication – Prescribing advice, interactions, availability of medicines**Evelina London Children’s Hospital Medicines Helpline | Tel: 020 7188 3003Email: Letstalkmedicines@gstt.nhs.uk  |
| **Guy’s and St. Thomas’ Hospital switchboard: 020 7188 7188** |
| **Medication – Prescribing advice, interactions, availability of medicines**Evelina London Children’s Hospital Medicines Helpline**GSTT Community Child and Adolescent Mental Health Service (CAMHS)**Dr Max DavieConsultant Community PaediatricianLambeth CAMHS 38 Black Prince Road | Tel: 020 7188 3003Email: Letstalkmedicines@gstt.nhs.ukTel: 020 3049 6004 Tel: 020 3228 7370 |
| **Lewisham and Greenwich Hospitals** **University Hospital Lewisham main switchboard: 020 8333 3000****Queen Elizabeth Hospital main switchboard: 020 8836 6000** |
| **Medication – Prescribing advice, interactions, availability of medicines**For University Hospital Lewisham patients please refer to SLAM details provided below.For Queen Elizabeth Hospital please refer to Oxleas NHS Trust details below |
| **South London and Maudsley (SLAM): 020 3228 6000** |
| **Medication – Prescribing advice, interactions, availability of medicines**Maudsley Medicines Information Services | Tel: 020 3228 2317  |
| **Oxleas NHS Trust switchboard 01322 625700** |
| Integrated Neurodevelopmental Team - ADHD Service  | Tel :020 8836 8621Email: oxl-tr.childrenstherapies@nhs.net |
| **Medication – Prescribing advice, interactions, availability of medicines**Oxleas Medicines Information | Tel: 013 2262 5002  |
| **Bromley Healthcare Community Services**  |
| **Consultant/specialist team**Dr Mark O'Leary,Consultant Community PaediatricianDenise Cox, ADHD Nurse  | Tel: 0208 466 9988 Email: Denise.Cox4@nhs.net Direct line: 0208 315 4757 |