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# A CONCISE REVIEW ON ANALYTICAL METHOD DEVELOPMENT AND VALIDATION OF OLANZAPINE

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## **Abstract:**

Olanzapine (OLZ) is an atypical antipsychotic agent is and different antipsychotic agent medications like Carbamazepine, Fluoxetine hydrochloride, Simvastatin, Clozapine, paliperidone, Quetiapine, several beta blocker, Risperidone, 9-Hydroxyrisiperidone, Demethylolanzapine, Aripiprazole, Orphenadrine, 1,2 Naphthoquinone, P-dimethylamino Benzaldehyde, Cerium sulphate, N-bromosulphinimide. The present investigation assesses the various approaches for analysis of OLZ in bulk drug as well as their pharmaceutical formulations. A concise survey states the collection and outline of about 74 explanatory strategies which incorporates HPLC, HPTLC, UV-Spectrophotometry, electrochemical techniques, LC-MS/MS, techniques actualized for examination of OLZ in biological matrices, bulk samples and in different dosage forms. The review depicts the rate usage of the different methodologies for examination of OLZ. The measurable information concerning the utility of these strategies for estimation of OLZ distributed during 1995 to 2018 have been incorporated.

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**Keywords:** Olanzapine; HPLC; HPTLC; LC-MS/MS; Spectrophotometry.

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# **INTRODUCTION:**

Olanzapine (OLZ) is an atypical antipsychotic agent and chemically it is (2-methyl-4-(4-methyl-1piperazinvl)-10H-thieno 3-b1 ſ2**.** [1, benzodiazepine) is an (figure 1).[1] It has antipsychotic activity and the usual dose is 10 mg once daily and it is a white to off-white powder OLZ It is used in the treatment of depression. [2]. Additionally, OLZ It decreases the gluconeogenesis while increasing the glucose uptake by muscles and fat cells. Olanzapine Works by blocking the receptors in the brain that are involved in transmitting these messages between the nerve cells. [3]. It has a higher affinity for 5-HT2 serotonin receptors than D2 dopamine receptors. OLZ is extensively metabolised to form 10-N-glucuronide, 4'-N-desmethyl, 2hydroxymethyl and 4'-N-oxide metabolites. Reacts with glucuronyltransferase to form a 10-Nglucuronide and a quaternary 4'-Nglucuronide. The cytochrome P450 (CYP) and (FMO) are responsible for OLZ metabolism. CYP1A2 and FMO3 form 4'-Ndesmethyl olanzapine and olanzapine Noxide, respectively (Figure 2). The average concentrations of olanzapine 10-N-glucuronide and N-desmethyl olanzapine. 2-Hydroxymethyl olanzapine is a minor metabolite and is primarily formed by CYP2D6. John T. Callaghan, Richard F. Bergstrom, Louis R. Ptak, and Charles M. Beasley

OLZ is also available in combination with The OLZ in various dosage forms as single constituent and in combination with Carbamazepine, Fluoxetine hydrochloride, Simvastatin, Clozapine, paliperidone, Quetiapine, several beta blocker, Risperidone, 9-

Hydroxyrisiperidone, Demethylolanzapine, Aripiprazole, Orphenadrine, 1,2 Naphthoquinone, Pdimethylamino Benzaldehyde, Cerium sulphate, Nbromosulphinimide. Although reviews about the pharmacology of OLZ have been earlier available, none of these reviews concentrated on OLZ analytical methods, perhaps because it is one of the bisystolic drug introduced in the market. The aim of this review is to deliver summary of the relevant published literature and a discussion of methods for the determination of OLZ on its own or in mixtures, in pure form, formulations, and biological samples using different analytical procedures (HPLC, HPTLC, UV, Bio analytical, LC-MS/MS, etc)

An extensive literature survey was done using the database like scholar, scifinder, Pubmed, Scopus and web of science. The literature survey revealed that the numerous analytical methods have been reported for OLZsuch as high-performance chromatography, high-performance thin-laver chromatography, and Liquid chromatography coupled with mass spectrophotometry, ultraviolet and visible spectrophotometry. [1-74]. Therefore, the aim of the proposed work to analysed and summarized all the analytical method exemplified in the literature. Taking into account of applicability of OLZ in the treatment of Antipsychotic. This analytical profile of OLZfocuses the analytical methods determination and quantification of OLZ pharmaceutical formulation as well as in biological samples as stated in the literature. Additionally, this review focuses on only the methods reported in the period of 1995-2018.

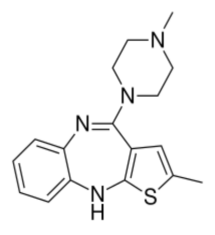


Figure 1. Chemical Structure of OLZ

## Metabolites of OLZ

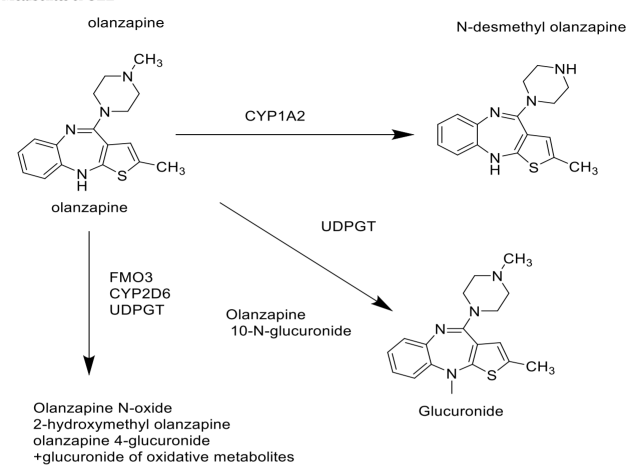


Figure 2. Metabolites of OLZ

## Pharmacopoeial status:

OLZ is the official drug in Indian Pharmacopoeia (IP) - 2007, Indian Pharmacopoeia (IP) - 2010, Indian Pharmacopoeia (IP) - 2014, the Merck index Thirteenth edition, the Merck index fourteen editions and Martindale.

IP depicted HPLC assay method using C18 (25 cm  $\times$  4.6 mm, 5 $\mu$ m) column as a stationary phase and mobile phase consisted of 3gm of ammonium dihydrogen orthophosphate adjust pH to 2.5, water and triethylamine (70:30  $\nu/\nu$ ) with a flow rate of 1 mL/min. Column effluent was monitored at 220 nm [5].

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rate of 1 mL/min. Column effluent was monitored at 220 nm [6].

IP depicted HPLC assay method using C18 (25 cm  $\times$  4.6 mm, 5 $\mu$ m) column as a stationary phase and mobile phase consisted of 4.83gm of sodium dihydrogen orthophosphate monohydrate adjust pH to 6.8, Acetonitrile and Methanol (80:20  $\nu/\nu$ ) with a flow rate of 1.2 mL/min. Column effluent was monitored at 230 nm [7].

BP depicted HPLC assay method using C18 (25 cm  $\times$  4.6 mm, 5 $\mu$ m) column as a stationary phase and mobile phase consisted of 0.345% w/v sodium dihydrogen phosphate monohydrate adjusted to pH 6.8 with dilute sodium hydroxide.buffer,Acetonitrile and Methanol (25:75 $\nu$ /v) with a flow rate of 2 mL/min. Column effluent was monitored at 250 nm [8].

# Analytical Methods for OLZ determination of HPLC Method:

Various separation techniques such HPLC have been employed for the determination of OLZ in pharmaceutical matrix as well as in biological samples. But, HPLC technique was mostly utilized for determination of OLZ. Most of the researchers utilized C18 reverse phase analytical column for separation of OLZ.

Maximum analytical studies with HPLC employed for determination of OLZ in pharmaceutical formulation on C18 reverse phase analytical columns. Water and Methanol in the ratio of 30:70 v/v or 70:30 v/v mixture as a mobile phase, associated or not with some organic additives such as buffers, acids or bases, methanol to improve selectivity and separation. The wavelength was used in HPLC methods ranges from 220-295 nm but in general it was set at 235 nm. The data related to simultaneous determination of drugs was specified in **Table 1** [9-18].

## **HPLC Simultaneous method development:**

Mahmoud A Tantawy et al. (2012) A spectrophotometry sensitive, accurate, & precise. TLC spectrodensitometry & HPLC method for simultaneous determined for OLZ & FLU-HCl. two

spectrophotometry techniques developed first derivative (D1) & derivative ratio (DD1). The precoated aluminium TLC plate with silica gel GF254 Stationary Phase & mobile phase mixture methanol: toluene: ammonia (7:3:0:1v/v/v) chromatogram wavelength at 235 nm.The HPLC developed methods used RP-C18 column with isocratic elution. The mobile phase mixture Acetonitrile: triethylamine (53:47:0.03v/v/v) pH adjusted 4 & flow Rate was found to be about 1.0ml/min. The wavelength at 235 nm. [11].

C. Vitorino et al. (2013) The RP-HPLC method developed by the Simultaneous determinate for SA, prodrug & the respective active hydroxy acid, SA & OZL in dosage form. Containing coencapsulating-Nanostructured lipid carries. The chromatography separation was carried by Phenomenex Luna phenylhexyl column, ( $5\mu m$ , $150\times 3$  mm) temperature at 35°C, & Socratic Conditions Used wavelength at 230 nm. The mobile phase mixture ammonium acetate aqueous solution 0.02M, methanol and Acetonitrile (30:35:35v/v/v) and flow rate was found to be about 0.8ml/min. The linear regression analysis, calibration curve good concentration range was found to be 0.5-100µg/mL, r2=0. 9994 for all three compounds, SA, OLZ. [12].

Table 1. Simultaneous estimation of drugs by HPLC

| Sr<br>no | Name of<br>drug/<br>formulati<br>on. | Column                                   | Mobile Phase<br>Composition                                                      | Detection (nm) Detector | Linearity<br>Retention<br>Time                                    | Flow rate (ml/min) | Ref |
|----------|--------------------------------------|------------------------------------------|----------------------------------------------------------------------------------|-------------------------|-------------------------------------------------------------------|--------------------|-----|
| 1        | OLZ+FL<br>U HCl<br>Tablet            | Inertsil<br>C18 ODS<br>column            | Acetonitrile: Methanol (90:10V/V)                                                | UV detector 233         | 20-80µg/ml<br>2.7min For<br>Fluoxentine<br>HCl and 3.3min         | 1                  | 9   |
| 2        | OLZ +<br>FLU HCl<br>Tablet           | C18<br>column                            | Acetonitrile: methanol:<br>0.032 M ammonium<br>acetate buffer<br>(45:05:50V/V/V) | UV/Visible detector 235 | 0.2-4μg/ml<br>0.1-2 μg/ml<br>300–1000 -<br>150–500 ng<br>1.95.min | 1.5                | 10  |
| 3        | OLZ +<br>FLU<br>Capsule              | Zorbax<br>ODS<br>column<br>C18<br>column | Phosphate buffer pH<br>4.0:acetonitrile:triethylam<br>inen(53:47:0.03V/V/V)      | UV detector 235         | 20–100 ug/mL.<br>100–600<br>ug/mL.<br>2.74, 9.77min.              | 1.0                | 11  |

| 4  | OLZ+SA                                          | Luna<br>Phenyl<br>Hexyl,c18<br>column | Ammonium acetate<br>aqueous solution<br>0.02Mmethanol:Acetonitr<br>ile (30:35:35 V/V/V)                                                    | UV/visible detector            | 0.5-100 μg/mL<br>7 min                                      | 0.8 | 12 |
|----|-------------------------------------------------|---------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------|-------------------------------------------------------------|-----|----|
| 5  | OLZ+CL<br>Z+QUE<br>Several<br>beta-<br>blockers | RP-4 ADS<br>column                    | Ethanol : water (80:20 V/V)                                                                                                                | PDA Detector 215               | 20-80 μg/ml<br>10.89 min                                    | 1   | 13 |
| 6  | OLZ+FL<br>U HCl<br>Tablet                       | C18<br>column                         | Acetonitrile: pot. Dihydrogen phosphate buffer: trietythimine (0.2%) (0.1% v/v ortho phosphoric acid PH3.1) (40:60:02V/V/V)                | UV detector 233                | 10 - 60μg/ml<br>20-120μg/ml<br>1.96&1.59min                 | 1.0 | 14 |
| 7  | OLZ<br>+FLU<br>Tablet                           | C18<br>column                         | 75mM potassium<br>dihydrogen phosphate<br>buffer (pH4.0):<br>Acetonitrile: methanol<br>(55:40:5V/V/V)                                      | UV /visible<br>detector<br>227 | 5–80μg/mL<br>& 20–320<br>μg/mL                              | 0.8 | 15 |
| 8  | OLZ+FL<br>U HCl<br>Tablet                       | HYPERSI<br>L ODS<br>C18<br>column     | 0.01M Phosphate buffer<br>PH 5.8: Acetonitrile<br>(55:45V/V) pH-2.6<br>adjusted with<br>Orthophosphoric acid)                              | PDA detector<br><b>261</b>     | 18-42µg/ml<br>and<br>72-168µg/ml.<br>3.480 and 2.597<br>min | 1   | 16 |
| 9  | OLZ+FL<br>U HCl<br>Tablet                       | C8<br>column                          | 0.1% v/v Ortho Phosphoric acid in water (pH 3.5 With Triethylamine):Acetonitril e: Methanol (60:30:10V/V/V)                                | PDA detector<br>225            | 12-28μg/ml<br>48-112μg/ml<br>2.19 min and<br>3.71.          | 1.0 | 17 |
| 10 | OLZ<br>Tablet                                   | Inertsil<br>C18<br>column             | 9.5 mM sodium<br>dihydrogen phosphate<br>(pH adjusted to 6.8 ± 0.1<br>with triethylamine) :<br>Acetonitrile : methanol<br>(40:30:30 V/V/V) | PDA detector 225               | 25 -75<br>μg/mL<br>100–300 μg/mL<br>10 min                  | 1.2 | 18 |

# HPLC method indicating stability study, impurity profiling methods

Ramisetti Nageswara Rao et al. (2008) The simultaneous determination of OLZ by RP- HPLC method. A process impurity in bulk drug & tablet dosage form was developed. The separation was

accomplished on Inertsil ODS 3V column (4.6mm, 250mm,  $5\mu$ m). The mobile phase mixture of 0.2M ammonium acetate (pH= 4.50): ACN in gradient elution mode. The analysis was PDA detector wavelength at 254 nm. The flow rate was found to be 1.0ml/min. [21].

Table 2. HPLC method indicating stability study, impurity profiling for OLZ

| Sr | Name of drug/ | Column | Mobile Phase | Detection | Linearity      | Flow     | Ref |
|----|---------------|--------|--------------|-----------|----------------|----------|-----|
| no | formulation.  |        | Composition  | (nm)      | Retention Time | rate     |     |
|    |               |        |              | Detector  |                | (ml/min) |     |

| 11 | OLZ Bulk               | Inertsil C18<br>column                                | Ammonium<br>phosphate buffer :<br>methanol (70:30<br>v/v)                              | UV-Visible detector SPD 10 220 | 2 - 10μg/ml<br>3.447min                                 | 1   | 19 |
|----|------------------------|-------------------------------------------------------|----------------------------------------------------------------------------------------|--------------------------------|---------------------------------------------------------|-----|----|
| 12 | OLZ+CMZP<br>Tablet     | ACE5-CN<br>column                                     | Phosphate buffer<br>(pH 5.0 25 mM):<br>methanol (80:20<br>v/v) (70:30 v/v)             | UV/DAD<br>detector<br>254      | 0.2–50.0 mg/mL                                          | 1   | 20 |
| 13 | OLZ<br>Tablet          | Inertsil<br>ODS 3V<br>column                          | Water: methanol (30:70v/v).                                                            | PDA detector 254               | 10 - 300 μg/mL                                          | 1.0 | 21 |
| 14 | OLZ<br>Tablet          | Intersil<br>ODS<br>column                             | Ammonium<br>acetate<br>(pH4.5):Acetonitr<br>ile<br>(70:30v/v)                          | PDA detector UV detector 271   | 10-200 mg/mL<br>7.48 min                                | 0.5 | 22 |
| 15 | OLZ<br>Tablet          | C18<br>column                                         | Potassium<br>dihydrogen<br>phosphate Buffer<br>(pH<br>6):Acetonitrile<br>(60:40) (v/v) | UV detector 258                | 5-25 μg/ml<br>5 min                                     | 1   | 23 |
| 16 | OLZ +PAL<br>Bulk       | C18, YMC<br>packpro<br>C18,<br>Inertsil<br>ODS 3V     | 0.1%Ammonium<br>Acetate in water :<br>Acetonitrile (95:5<br>v/v)                       | PDA detector 254               | 0.2 mg/ml and<br>0.5mg/ml<br>0.2 mg/ml and<br>0.5 mg/ml | 0.8 | 24 |
| 17 | OLZ<br>Bulk            | Agilent<br>Octyldecyl<br>silica<br>column<br>(TC-C18, | Methanol: 0.3%<br>TEA in water<br>(36: 64 v/v)                                         | UV/Visible detector 254        | 50 mg ml - 320<br>mg/ml<br>11.10 time /min              | 1.0 | 25 |
| 18 | OLZ<br>Tablets         | Kromasil<br>C-18<br>column                            | Acetonitrile: phosphate buffer (30:70 v/v)                                             | DAD<br>258                     | 10 - 50 μg /mL<br>1.850 min                             | 1.5 | 26 |
| 19 | OLZ<br>Tablet          | BDS<br>Hypersil<br>C18<br>Column                      | 0.01M Tetra butyl<br>ammonium<br>hydrogen<br>sulphate:<br>Methanol (60:40<br>v/v).     | UV Detector 228                | 10-80μg/ml<br>10.0 min.                                 | 1.0 | 27 |
| 20 | OLZ Bulk and<br>Tablet | Intersil<br>ODS 3V<br>column                          | 10mM disodium<br>hydrogen<br>phosphate buffer<br>(pH 7.4):<br>Acetonitrile (35:        | UV/ visible detector 254       | 2.5–20.0 µg/mL<br>4.39 ± 0.01min                        | 1.0 | 28 |

|    |            |         | 65v/v)            |     |              |     |    |   |
|----|------------|---------|-------------------|-----|--------------|-----|----|---|
| 21 | OLZ Tablet | ODS A-  | phosphate buffer  | DAD | 1.61×106 -   | 1.3 | 29 | l |
|    |            | 132 C18 | (pH: 5.5):        | 295 | 7.24×104 min |     |    | İ |
|    |            | column  | Acetonitrile (7:3 |     |              |     |    |   |
|    |            |         | v/v)              |     |              |     |    |   |

# **Bio-analytical Method:**

# The Bioanalytical Method development in Table 3

Christoph Hiemke et al. (2001) A simultaneous technique of the antipsychotic drug CLZ, OLZ, & demethylated metabolites. Method included adsorption CPS coated clean up column washes & interfering serum constituents to waste separation on ODS Hypersil C18 column RP- material (5 μm, 250 × 4.6 mm). A used mobile phase mixture of Acetonitrile: water: tetra methyl ethylene diamine (37:62.6.4v/v/v) pH adjusted 6.5 concentrated acetic acid. The UV detection Lamda max at 254 nm. The LOQ was found to be 10-20ng/ml. Relative standard variation ranges between 4.5 and 13.5 [33].

Huande Li et al. (2012) sensitive, rapid LC-MS/MS technique coupled with column developed for the determined of OLZ in rat brain microdialysates. The both columns C8 guard column used samples before analysis separate on a C18 column & detection with tandem mass spectrometry. The both mobile phase mixtures of methanol: Acetonitrile: water

(43:43:14v/v/v) was for analysis separated, water in both mobile phases contained 0.1% ammonium acetate. The LOQ for OLZ was found to be 0.085ng/ml. The linear from LOQ to 34ng/ml with a coefficient of determination more than 0.998. Precision study Intraday and interday& accuracy were determined with variability less than 13.24% (RSD). [35].

M. a. raggi et al. (2001) The HPLC method with electrochemical detection has been developed for the determination of olanzapine and its main metabolite, desmethylolanzapine, in human plasma. Chromatographies separation and analysis was performed on a C8 reversed phase column with a mixture of methanol. Acetonitrile, and pH 3.7 buffer phosphate as mobile phase, methylolanzapine was used as internal standard. The response was linearly dependent on concentration and precision were satisfactory over the concentration range 0.5-75.0ng/ml for both analytes. The limit of detection was 0.2ng/ml for both analytes. [36].

Table 3. Bioanalytical paper

| Sr<br>no | Name of<br>drug        | Sample<br>matrix | Column                        | Mobile phase composition                                                                                                       | Detection (nm) detector       | Flow<br>rate<br>(ml/min) | Ref | Internal<br>standard |
|----------|------------------------|------------------|-------------------------------|--------------------------------------------------------------------------------------------------------------------------------|-------------------------------|--------------------------|-----|----------------------|
| 1        | OLZ+ RIS+<br>9-HYD RIS | Human<br>Plasma  | RP18<br>column                | 10 mM<br>ammonium acetate<br>buffer at a pH of<br>3.5 which was<br>adjusted with<br>acetic acid:<br>Acetonitrile<br>(70:30v/v) | PDA detector 277              | 0.3                      | 30  | Clozapine            |
| 2        | OLZ                    | Human<br>Plasma  | YMC<br>column                 | 75 mM sodium phosphate (pH 7): methanol: Acetonitrile (48:26:26 v/v/v).                                                        | Electrochemic<br>al detection | 1.2                      | 31  | Olanzapin<br>e       |
| 3        | OLZ                    | Rat Plasma       | column<br>C18hypers<br>il-BDS | 50 mM phosphate<br>buffer pH 5.5:<br>Acetonitrile :<br>methanol                                                                | UV detector                   | 1.2                      | 32  | Olanzapin<br>e       |

|    |                         |                              |                                          | (50:30:20 v/v/v)                                                                                              |                              |      |    |                                               |
|----|-------------------------|------------------------------|------------------------------------------|---------------------------------------------------------------------------------------------------------------|------------------------------|------|----|-----------------------------------------------|
| 4  | OLZ                     | Human<br>Breast Milk         | YMC<br>basic<br>column                   | 75mM phosphate<br>buffer pH 7.0:<br>acetonitrile:<br>methanol<br>(48:26:26v/v/v)                              | Electrochemic al detection   | 1    | 33 | Olanzapin<br>e                                |
| 5  | OLZ                     | Serum                        | Normal-<br>phase<br>silica gel<br>column | 50 mM<br>ammonium acetate<br>buffer adjusted to<br>pH 9.9 with<br>ammonia water:<br>methanol<br>(15:85v/v).   | UV detector 270              | 1.1  | 34 | Trifluoper<br>azine                           |
| 6  | OLZ+CLZ                 | Serum                        | C18 ODS<br>Hypersil                      | Acetonitrile: water tetramethyl ethylene diamine(37:62.6:0. 4 v/v/v)                                          | UV detector 254              | 1.5  | 35 | Olanzapin<br>e and<br>demethylo<br>lanzapine. |
| 7  | OLZ+CLZ<br>+RISP<br>QUT | Rat Brain                    | Macherey<br>nagel C18<br>column          | water (formic acid: 2.70 mmol/l ammonium acetate: 10 mmol/l): Acetonitrile (53:47v/v)                         | UV detector 254              | 0.16 | 36 | Olanzapin<br>e and<br>quetiapine              |
| 8  | OLZ                     | Human<br>plasma              | C18<br>column                            | 0.06 M<br>ammonium<br>acetate buffer pH<br>5.9 : Acetonitrile :<br>methanol (40: 41<br>:3 : 7 v/v/v)          | electrochemic<br>al detector | 0.69 | 37 | Clozapine                                     |
| 9  | OLZ                     | Human<br>plasma and<br>Urine | Supelcosil<br>LC-CN<br>column            | 10% methanol25%<br>acetonitrile and<br>65% 50 mM<br>phosphate buffer<br>pH 6.0                                | UV detector 214              | 1    | 38 | Olanzapin<br>e                                |
| 10 | OLZ                     | Rat plasma                   | YMC<br>basic<br>column                   | 75mM phosphate buffer (adjusted to pH 7 with 5 M  Sodium hydroxide): Acetonitrile: methanol  (48:26:26 v/v/v) | electrochemic<br>al detector | 1.2  | 39 | Olanzapin<br>e                                |

| 11 | OLZ             | Human<br>Plasma | Spherisorb<br>S5 C6<br>analytical<br>column | Water:<br>Acetonitrile<br>(55:45 v/v)                                                                                                 | UV-VIS<br>detector<br>254         | 1.0 | 40 | Clozapine                    |
|----|-----------------|-----------------|---------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------|-----|----|------------------------------|
| 12 | OLZ             | Human<br>Plasma | C 18<br>column                              | 14% acetonitrile in<br>water (containing<br>0.25% H PO and<br>0.05%<br>triethylamine)                                                 | electrochemic al detectors 270    | 1   | 41 | N-<br>desmethyl<br>clozapine |
| 13 | OLZ+DES<br>MOLZ | Human<br>Plasma | C8<br>column                                | Methanol(11%o),a<br>cetonitrile(9.7%),<br>and 8.9mmol L 1<br>phosphate<br>buffer(79.3%)cont<br>aining 7.18mmolL<br>1<br>Triethylamine | Decade amperometric detectors 316 | 0.7 | 42 | Methylola<br>nzapine         |
| 14 | OLZ             | Human<br>Plasma | column C8                                   | Acetonitrile:13.5m<br>M, pH 2.0<br>phosphate buffer<br>(30:70 v/v pH =<br>2.5)                                                        | UV detector 260                   | 1   | 43 | Tripralidin<br>e             |
| 15 | OLZ+ARI         | Human<br>plasma | column<br>Rp-18                             | Phosphate buffer (pH 3.14, 20 mM) and acetonitrile                                                                                    | Diode array detector. 255         | 0.8 | 44 | Carbamaz<br>epine            |

# **Spectrophotometric method:**

Spectrophotometric methods with UV-Visible detection have been developed for OLZ analysis without combination illustrated in Table 4, 5, 6 [45-62]

Kishanta Kumar Pradhan et al. (2014) The Olanzapine in pure and tablet form. The simple, specific and reliable UV-VIS spectrophotometry method was studied. The bulk forms mobile phase mixture of water: hydrochloric acid (9:1). The Lamda max at 258 nm. The regression equation and calibration graph, respectively. The concentration range of  $5\text{-}40\mu\text{g/ml}$ . The correlation coefficient was found = 0.059x + 0.171 and 0.998 respectively [45].

S.firdous et al. (2005) The determination of olanzapine, based on UV spectrophotometry new method and non-aqueous titration, has been developed. The Lamda max at 226 nm. In a

methanolic solution of olanzapine. The concentration range was found to be 0.1g/ml to 50g/ml beers law obeys and interday precision of UV is 0.97%. The non-aqueous titration was carried by olanzapine with 0.1N Perchloricacid. Using naphthobenzene as indicators and Intraday precision ranges 0.35%. [50].

Sahar R. Fadhel et al. (2016) The new spectrophotometric techniques have been developed for the assay of olanzapine in bulk and tablet forms. This both methods, in acidic medium based on the diazocouping of olanzapine and diazotized p-Nitroaniline to form a stable brown colored water-soluble azo dye. The maximum Lamda max at 405 nm. The concentration linear range was found to be  $0.5\text{-}45.0~\mu\text{g/ml}$ .  $1.5777\times104~\text{L}$ . Mol-1. Cm the LOD was found to be  $0.3148~\mu\text{g}$ . mL-1 and sandals sensitivity values were found to be  $0.0198~\mu\text{g/cm}$ . [53].

Table 4. UV- Spectroscopy method

| Sr.no | Drug | Method<br>Wavelength<br>(nm)                         | Matrix                                   | Linearity/lod/loq                                                                                 | Ref |
|-------|------|------------------------------------------------------|------------------------------------------|---------------------------------------------------------------------------------------------------|-----|
| 1     | OLZ  | UV Spectroscopic<br>Method<br>258                    | Bulk                                     | 5-40μg/ml<br>LOD=0.4306ug/ml<br>LOQ=1.305ug/ml                                                    | 45  |
| 2     | OLZ  | Spectrophotometric methods 410 ,620                  | Balk Formulation Using Bromocresol Green | 0.25-12.5µg/ml and 0.2-<br>5.0ug/ml<br>LOD=0.28and 0.03ug/ml<br>LOQ=0.86 and 0.08ug/ml            | 46  |
| 3     | OLZ  | UV Spectrophotometric Methods 258,252                | Bulk                                     | 3-18 µg/ml<br>and 4-24 µg/ml<br>LOD=0.1680and<br>0.2018ug/ml<br>LOQ=0.5091and<br>0.6115ug/ml      | 47  |
| 4     | OLZ  | Spectrophotometry Methods 550, 610                   | Tablets                                  | 2.0 -20μg/ml<br>and 1.0- 10 μg mL<br>LOD and LOQ=0.37 and<br>1.13 μg mL<br>0.16 And 0.48 μg mL-1. | 48  |
| 5     | OLZ  | UV<br>Spectrophotometric<br>method<br>270, 304 & 304 | Tablets                                  | 10-16 μg/ml<br>LOD=0.101,<br>0.209,0.109ug/ml<br>LOQ=0.306,<br>0.634,0.332ug/ml                   | 49  |
| 6     | OLZ  | UV Spectrophotometry and non-aqueous titration 226   | Tablet                                   | 0.1 - 50ug/ml<br>0.2 - 100mg/m<br>LOD=0.1ug/ml,0.3mg                                              | 50  |
| 7     | OLZ  | Spectrophotometric methods 222, 230                  | Bulk                                     | 2-10 µg/ml<br>LOD=500ug/ml &<br>499ug/ml<br>LOQ=166.6ug/ml &<br>159.2ug/ml                        | 51  |
| 8     | OLZ  | Highly sensitive Spectrophotometric method 610       | Bulk                                     | 0.3 - 8.0ug/ml<br>LOD=0.1290ug/ml<br>LOQ=0.3696ug/ml                                              | 52  |
| 9     | OLZ  | Spectrophotometric method 405                        | Bulk                                     | 0.4 - 45ug/ml<br>LOD=0.3148ug/ml<br>LOQ=1.0495ug/ml                                               | 53  |

# Simultaneous method development

Farzana I Ghanchivhora et al. (2017) The spectrophotometry techniques simple, specific, accurate, precise and economical has been developed for the both drug olanzapine and paliperidone in

synthetic mixture. The olanzapine maximum Lamda max at 259 NM & paliperidone maximum Lamda max at 269 nm. The concentration liner range of olanzapine 2-12  $\mu$ g/ml and paliperidone range 3-18  $\mu$ g/ml [57].

Table 5. Simultaneous UV- Spectroscopy method

| Sr.<br>no | Drug                                       | Method<br>Wavelength<br>(nm)                         | Matrix | Linearity<br>Lod/Loq                                                       | Ref |
|-----------|--------------------------------------------|------------------------------------------------------|--------|----------------------------------------------------------------------------|-----|
| 10        | OLZ+FLU HCI                                | Simultaneous<br>Methods<br>318,239                   | Bulk   | 10 - 60 mg/ml<br>LOQ= 0.73 to 1.49 mg/ ml and<br>0.18 to 0.96 mg/ ml       | 54  |
| 11        | OLZ+CLOZ+<br>QUES+several<br>beta-blockers | Simultaneous<br>Methods<br>215, 226, 242<br>and 299. | Tablet | 20-80 μg/ml<br>LOQ=2.5μg/ml<br>LOD=2.50 μg/ml                              | 55  |
| 12        | OLZ+FLU HCl                                | Simultaneous method 226,258                          | Bulk   | 10-100 μg/ml<br>10-100 μg/ml<br>LOD=1-10 μg/ml<br>LOQ=10-50 μg/ml          | 56  |
| 13        | OLZ+ PAL                                   | Simultaneous method 269, 259                         | Bulk   | 3-18 μg/ml and 2-12 μg/ml<br>LOD=0.2131 0.645μg/ml<br>LOQ=0.218 0.662μg/ml | 57  |

# Analysis with combination drug

HD Revanasiddappa et al. (2012) The spectrophotometry method simple, sensitivity determined by OLZ &ORPDN bulk dosage form. The method developed is based on the ternary complex formulation of drugs under investigation with Essen and lead (II) by using methyl cellulose as a surfactant. The maximum wavelength at 540 nm both drug OLZ & ORPDN. The optimum experimental conditions for the ternary complex formulation established. The both methods obeys bees law concentration range was found to be 0.0-35.0 and 0.0-55  $\mu$ g/ml. [58].

**Table 6.** UV – Spectroscopy with combination drug

| Sr.no | Drug                                             | Method<br>Wavelength<br>(nm)            | Matrix  | Linearity/lod/loq                                                                                                 | Ref |
|-------|--------------------------------------------------|-----------------------------------------|---------|-------------------------------------------------------------------------------------------------------------------|-----|
| 14    | OLZ +ORPHE                                       | Sensitive Spectrophotometric Method 540 | Bulk    | 0.0-35.0μg/ml and 0.0-55<br>μg/mL<br>LOD=0.4547 and<br>0.9422μg/ml<br>LOQ= 0.1501 and<br>0.3109 μg/ml             | 58  |
| 15    | OLZ+ 1,2<br>NAPTHOQUI<br>-4 SUL (NQS)            | Spectrophotometric method 454           | Tablet  | 0.4 - 4.0μg/ml<br>LOD=0.09μg/ml<br>LOQ=0.29μg/ml                                                                  | 59  |
| 16    | OLZ +P-<br>DIMETHYL<br>AMINO<br>BENZ             | Spectrophotometric method 410           | Bulk    | 5–160μg/ml<br>LOD=6.6μg/ml<br>LOQ=20μg/ml                                                                         | 60  |
| 17    | OLZ +CER<br>(IV) SUL                             | Visible Spectrophotometry 480, 640, 700 | Tablets | 0.2-2.0μg/ml0.5-9.0μg/ml<br>0.2- 3.0μg/ml<br>LOD=0.01, 0.04, 0.01μg/ml<br>LOQ=0.02, 0.11,<br>0.03μg/ml            | 61  |
| 18    | OLZ+ N<br>BROMO<br>SUSSIMIDE<br>+CERB(IV)S<br>UL | Spectrophotometric method 532, 538 ,538 | Bulk    | 10 – 120μg/ml 0.5 – 6.0μg/ml 0.6 – 3.0μg/ml  LOD=2.10ug/ml 0.10μg/ml 0.16μg/ml  LOQ=6.99μg/ml 0.30μg/ml 0.37μg/ml | 62  |

**High-performance thin layer chromatography (HPTLC):** 

The HPTLC technique overcomes the limits of TLC as well as takes advantage over the HPLC techniques. It shows the advantages like less time consuming,

cheap, utilizes disposable stationary phases, less sample required compared to TLC, gives static and offline detection ability and high throughput qualitative and quantitative detection. The analytical information about HPTLC illustrated in **Table 7** [63-66].

Sejal Patel et al. (2009) The two different techniques a binary mixture of fluoxentine Hcl and olanzapine. The first method RP-HPLC determined of fluoxentine Hcl and olanzapine. The mobile phase mixture of Acetonitrile: methanol: 0.032 M ammonium acetate buffer (45:05:50, v/v/v). The flow rate was found to be 1.5ml/min. Lambda max at 235

nm. The concentration range was found to be 0.2- $4\mu g/ml$  and 0.1- $2\mu g/ml$ . The both drug % recovery was found to be  $101.16\pm0.59$  and  $99.79\pm0.56\%$ . The second method HPTLC the both drugs, separation followed by densitometry measured of their spots at 235nm. The separation carried out by Merck TLC aluminium plate of silica gel 60F254. The mobile phase mixture of acetone: methanol: triethyleamine (5:3:0:5v/v/v). The linearity was found to be in the range of 300-1000ng/spot and 150-500ng/spot and % recovery were found to be  $100.95\pm0.52$  and  $99.31\pm0.51\%$  for Fluoxetine HCl and olanzapine. [65].

**Table 7.** HPTLC determination of OLZ

| Sr<br>no | Drug                                               | Formulat<br>ion | Stationary Phase Plate   | Mobile Phase<br>Composition                                                                   | Detection in (nm) | linearity                                      | Rf                                                  | Ref |
|----------|----------------------------------------------------|-----------------|--------------------------|-----------------------------------------------------------------------------------------------|-------------------|------------------------------------------------|-----------------------------------------------------|-----|
| 1        | OLZ+<br>FLU                                        | Tablet          | Silica gel<br>60F254     | Methanol:<br>toluene<br>(4:2v/v)                                                              | 233               | 100-800<br>ng/spot<br>1000-8000<br>ng/spot     | 0.31±0<br>.01                                       | 63  |
| 2        | OLZ+<br>DUOH<br>Cl<br>Synthe<br>tic<br>Mixtur<br>e | Capsule         | Silica<br>gel 60<br>F254 | Toluene: methanol:10% ammonia3:1.3:0 .05 (v/v)acetone: methanol: triethylamine(5: 3:0.5v/v/v) | 231,<br>240       | 60–480 ng/spot<br>per<br>100-800&<br>50–400 ng | 0.39<br>±0.02<br>0.63<br>±0.02<br>& 0.77<br>± 0.02, | 64  |
| 3        | OLZ+<br>FLU<br>HCl                                 | Tablet          | Silica gel<br>60 F254    | Acetone:methan ol:triethyleamin e (5:3:0.5 v/v/v)                                             | 235               | 300–1000 &<br>150–500 ng/spot                  | -                                                   | 65  |
| 4        | OLZ                                                | Bulk            | Silica gel<br>60F254     | Methanol :ethyl<br>acetate (8.0 +<br>2.0 v/v)                                                 | 285               | 100 - 600<br>ng/band                           | Rf = 0.35 - 0.02                                    | 66  |

## LC-MS/MS method:

Coupling of HPLC with single MS or MS-MS is highly sensitive, able to analyze multicomponent, to inspect the specificity of the analysis and most reliable method to evaluate active ingredients from

biological samples. In existing LC/MS/MS techniques, ionization of the sample is carried out under atmospheric pressure (atmospheric pressure ionization (API) which is separated from the high vacuum portion of the mass analyzer. Two commonly

used methods of atmospheric pressure ionization involve electrospray ionization (ESI) and atmospheric pressure chemical ionization (APCI) of the molecules to be analyzed. The most use mobile phase was 10mM aqueous ammonium acetate adjusted to pH 4 with formic acid and Acetonitrile along with methanol. And for analysis of sample generally used matrix was Human plasma. The mass to charge ratio was used in the range of 313.2-256.3 and extraction of samples done by various methods such as protein precipitation, liquid-liquid extraction, solid-liquid extraction, Serum or cerebral spinal fluid samples, ion-exchange cartridges, Rat brain homogenate and analyzed etc. The separation was carried out on the

C18 column generally. The concise analytical data related to the LC-MS/MS is depicted in **Table 8 [67-82].** 

Michael G. Bartlett et al. (2007) The analytical method extract from rat brain homogenate and analyzed by LC-MS/MS. sample prepared and chromatography study, he method used a Water Atlantis TM dC-18 column (30mm×2.1mm 3mm). Gradient elution the mobile phase mixture of Acetonitrile: 5mM ammonium Formate adjusted pH6.1 with formic acid. The analytical method in positive ion separated used multiple reaction monitoring. [68].

Table 8, LC-MS

| Sr. | Drug                                                             | Matrix              | Extraction method                         | m/z ratio                                            |                                  | LC Separation                                                                                                                                                                                                | Lod/Loq                     | Ref |
|-----|------------------------------------------------------------------|---------------------|-------------------------------------------|------------------------------------------------------|----------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------|-----|
| no  |                                                                  |                     | method                                    |                                                      | Column                           | M.P                                                                                                                                                                                                          |                             |     |
| 1   | OLZ                                                              | Human<br>Plasma     | solid phase<br>extraction                 | m/z<br>313/256<br>m/z<br>384/253                     | ACE5<br>C18-300<br>column        | (5:95,10:90,15:85,20:80 and 30:70v/v) of water methanol: Acetonitrile: formic acid: ammonia(0.010.005%) ammonium trifluoroacetate ammonium acetate or ammonium formate buffer sin varying strengths (2–20mM) | LOD=0.10&0<br>.012<br>ng/mL | 67  |
| 2   | OLZ+RI<br>SP,9<br>HYDR<br>ORISP+<br>CLOZ+<br>HOPO<br>+ZIPRA<br>S | Rat Brain<br>Tissue | Rat brain<br>homogenae<br>and<br>analyzed | m/z 313m/z<br>256 32V 23<br>eV                       | C8 guard column.                 | 100µl of methanol: 20mM ammonium formate (pH3.86, adjusted by formic acid) (70:30v/v).                                                                                                                       | LOQ=0.208<br>ng/ml          | 68  |
| 3   | OLZ                                                              | Human<br>Blood      | Liquid—<br>liquid<br>extraction           | ( <i>m</i> / <i>z</i> 313.42<br>56.2)327.3<br>270.1) | Monochr<br>m HPLC<br>column      | 100 mM ammonium acetate: methanol: isopropanol: water (15:4:1v/v/v).                                                                                                                                         | -                           | 69  |
| 4   | OLZ+F<br>LU                                                      | Human<br>Plasma     | Plasma<br>samples on<br>Waters            | m/z 313,<br>310,<br>316and 315                       | Themo<br>Hypersil<br>Gold<br>C18 | 2 mM ammonium acetate:<br>Methanol (10:90 v/v)                                                                                                                                                               | LOQ=0.50<br>ng/ml           | 70  |

| 5  | OLZ                                                         | Rabbit<br>Plasma                                    | Liquid-<br>Liquid<br>extraction                 | m/z 313.4<br>→256.3                                               | C18<br>column                                         | 0.1% v/v formic acid in water<br>: Methanol (08:92 v/v)                                                                | LOQ=5ng/ml                                                                                                  | 71 |
|----|-------------------------------------------------------------|-----------------------------------------------------|-------------------------------------------------|-------------------------------------------------------------------|-------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------|----|
| 6  | OLZ+F<br>LU                                                 | Human<br>Plasma                                     | Solid phase extract on                          | m/z<br>310.01!147<br>.69<br>313.15!256<br>.14<br>298.1!153.<br>97 | Gold<br>C18colu<br>mn                                 | Acetonitrile: water containing 2% formic acid (70:30v/v)                                                               | LOQ=0.37<br>ng/ml                                                                                           | 72 |
| 7  | OLZ                                                         | Human<br>Plasma                                     | Liquid—<br>liquid and<br>SPE<br>extraction      | m/z 313.3<br>→ 256.1                                              | YMC-<br>ODS-AQ<br>C18<br>Column                       | 10 mM ammonium acetate in water contained 0.05% (v/v) formic acid (pH 3.5) methanol containing 0.05% (v/v) formic      | LOQ=0.2<br>ng/ml                                                                                            | 73 |
| 8  | OLZ                                                         | Human<br>Plasma                                     | Liquid-<br>liquid<br>extraction                 | m/z 313.15                                                        | ZORBA<br>XEclipse<br>XDB-CN<br>column                 | Acetonitrile: aqueous ammonium acetate solution (pH 4.0, 10 mM) (56:44v/v)                                             | LOQ=0.5ng/<br>mL                                                                                            | 74 |
| 9  | OLZ                                                         | Blood                                               | Solid-phase extraction                          | $m/z 313 \rightarrow 256$ $m/z 313 \rightarrow 84$                | Revered<br>phase<br>Zorbx<br>Extend-<br>C18colu<br>mn | Methanol : Acetonitrile : ammonium hydroxide (25:25:50 : 5 mM v/v/v)                                                   | LOQ=0.005m<br>g/kg                                                                                          | 75 |
| 10 | OLZ+RI<br>SP+QU<br>E+CLO<br>Z+ZIPR<br>AS+PE<br>RO+<br>ARIPI | Human<br>Serum                                      | Solid-phase extraction                          | -                                                                 | Mightysi<br>1-RP-18<br>MS<br>column                   | 10 mM formic ammonium<br>buffer (pH 6.0) : Acetonitrile                                                                | LOD=0.0007<br>1, 0.031,<br>0.015,<br>0.046, 0.017,<br>0.0057, 0.012<br>and<br>0.027ng/ml<br>LOQ=20ng/<br>ml | 76 |
| 11 | OLZ+N - DESME THYL OLZ                                      | Human<br>Serum<br>and<br>Cerebros<br>pinal<br>Fluid | Serum or<br>cerebral<br>spinal fluid<br>samples | -                                                                 | Hydro-<br>RP<br>column                                | buffer (10mM 0.05% formic<br>acid dilution of 5ml<br>ammonium formate stock<br>Solution and 250ml formic<br>acid (98%) | LOQ=0.3<br>ng/ml& 0.9<br>ng/ml                                                                              | 77 |

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| 12 | OLZ+D<br>ESMET<br>HYL<br>OLZ | Anticoag<br>ulant and<br>lipemia | Waters Oasis MCX cartridges and analyzed Solid-phase | m/z<br>312.9/256.<br>0                                                    | Phenome<br>nex<br>LUNA<br>pheyl<br>hexyl,col<br>mn | Acetonitrile: ammonium acetate (20 mM) (52:48v/v).  Formic acid : Acetonitrile (0.1:100 v/v). | -                 | 78 |
|----|------------------------------|----------------------------------|------------------------------------------------------|---------------------------------------------------------------------------|----------------------------------------------------|-----------------------------------------------------------------------------------------------|-------------------|----|
| 13 |                              | Urine                            | extraction                                           | -                                                                         | column                                             | Ammonium acetate (pH 7.8):<br>Acetonitrile (10:90v/v).                                        | LOQ=<br>1ng/ml    | 79 |
| 14 | OLZ                          | Human<br>Plasma                  | Liquid—<br>liquid<br>extraction.                     | m/z: 313.1<br>> 256.1<br>278.1 ><br>260.2                                 | ACE<br>C18,<br>column                              | water with 0.1% formic acid<br>Acetonitrile: 0.1% formic<br>acid (50: 50 v/v)                 | LOQ=<br>1ng/ml    | 80 |
| 15 | OLZ+F<br>LU+NO<br>R FLU      | Human<br>plasma                  | Liquid-<br>liquid<br>extraction                      | m/z 313.10<br>→256.05,<br>m/z 310.10<br>→148.00,<br>m/z 296.05<br>→133.90 | Agilent<br>Eclipse<br>Plus C18<br>column           | Methanol: 20 mM<br>ammonium formate buffer<br>(82.5: 17.5 v/v).                               | LOQ=0.05<br>ng/ml | 81 |
| 16 | OLZ                          | Human<br>plasma                  | Liquid—<br>liquid<br>extraction                      | m/z<br>313/256                                                            | Revere<br>phase<br>C18<br>column                   | 10mM ammonium acetate<br>buffer: Acetonitrile<br>(10:90v/v)                                   | LOQ= 100<br>pg/ml | 82 |

# **CONCLUSION:**

The present review illustrates various analytical approaches executed for the valuation of OLZ. An abundant investigation had performed, including, HPLC, Bio-analytical, HPTLC, UV/Vis-Spectroscopy, LC-MS/MS, GC-MS, etc. for estimation of OLZ in bulk and in its combined pharmaceutical formulations and in plasma. High performance-Liquid chromatography detection has been found to be the most studied for estimation of OLZ in bulk as well as pharmaceutical formulations, while hyphenated LS-MS/MS method was reported for determination of OLZ and its

metabolite in plasma and other biological fluids. Further, methods were reported for its pharmacokinetics as well as bioequivalence studies. Few chromatography approaches like HPTLC and Stability-indicating HPLC analysis is also reported in the literature. Certain Spectrophometric methods in UV-Visible spectroscopy analysis is most often used for assessment for OLZ.

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## **Conflict Of Interest:**

Authors do not have conflict of interest for this manuscript.

## **Abbreviations:**

- OLZ Olanzapine
- ♣ FLU- Fluoxetine
- CLZ- Clozapine
- ♣ CMZP- Carbamazepine
- PAL- Paliperidone
- ♣ RIS- Risperidone
- ♣ ARI- Aripiprazole
- ORPHE- Orphenadrine
- **♣** SA- Simvastatin
- LC-ES/MS/MS- Liquid chromatography Electrospray-mass spectroscopy-mass spectroscopy
- GC-MS-MS- Gas chromatography- mass spectroscopy-mass spectroscopy
- LC-MS- Liquid chromatography-mass spectroscopy
- **♣** SEM- Simultaneous equation method
- RF- Retention factor
- ♣ ESI- Electro-spray ionization
- nm-Nanometer
- M.P.- Melting point
- ♣ ACM-Absorption correction method
- ♣ ACN- Acetonitrile
- ♣ FA- Formic acid
- ♣ MFE- Mercury film electrode
- ♣ HMDE- Hanging mercury drop electrode
- **↓** CZE- Capillary zone electrophoretic
- MEKC- Micellar electro kinetic capillary chromatographic

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